

GenCore version 5.1.6  
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OM protein - protein search, using sw model

Run on: October 19, 2004, 18:56:12 ; Search time 158 Seconds

(without alignments)  
43.138 Million cell updates/sec

Title: US-10-799-005A-1

Perfect score: 97

Sequence: 1 EPNHLNGKIAFKIVSQBP 19

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2002273 seqs, 358729299 residues

Total number of hits satisfying chosen parameters: 249703

Minimum DB seq length: 19  
Maximum DB seq length: 50

Post-processing: Minimum Match 0%  
Maximum Match 100%  
Listing first 500 summaries

Database : A\_Geneseq\_23Sep04:\*

1: Geneseq1980s:\*

2: Geneseq1990s:\*

3: Geneseq2000s:\*

4: Geneseq2001s:\*

5: Geneseq2002s:\*

6: Geneseq2003as:\*

7: Geneseq2003bs:\*

8: Geneseq2004s:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match %	Length	DB ID	Description
1	38	39.2	22	2	AAR43461 Ro/SSA ep
2	35	36.1	46	3	AB27169 Sendai vi
3	34	35.1	45	6	ABM72186 Staphyloc
4	34	35.1	46	3	AB27167 HPIV3 par
5	34	35.1	49	4	AU21220 Human nov
6	33	34.0	26	4	AAM33005 Peptide #
7	33	34.0	26	4	AAM72776 Human bon
8	33	34.0	26	4	AAM60161 Human liv
9	33	34.0	26	4	ABG54477 Human liv
10	33	34.0	26	5	ABG42601 Human pep
11	33	34.0	44	3	AG27280 Zee may
12	32.5	33.5	36	3	AG59652 Arabidops
13	32.5	33.5	36	4	AAM16072 Peptide #
14	32.5	33.5	36	4	AB335064 Peptide #
15	32.5	33.5	36	4	AAM28566 Peptide #
16	32.5	33.5	36	4	AB20476 Protein #
17	32.5	33.5	36	4	AAM68248 Human bon
18	32.5	33.5	36	4	AAM5878 Human bra
19	32.5	33.5	36	4	ABG49902 Human liv
20	32.5	33.5	36	4	AAM037799 Peptide #
21	32.5	33.5	36	5	ABG37783 Human pep
22	32	33.0	24	6	AB012904 Mouse zin
23	32	33.0	24	6	AB011938 Human zin
24	32	33.0	24	6	ABU62168 Human zin
25	32	33.0	26	2	AAM60192 Bacteriop

99	30	30.9	39	6	ADA03571	Insulin r	172	29	29.9	41	8	ABO59597	Abos9597 Human gen
100	30	30.9	39	6	ABO34599	Region of	173	29	29.9	42	4	ABAB64605	Abab64605 Human sec
101	30	30.9	39	7	ADH94784	Insulin r	174	29	29.9	43	3	AAG09091	Agag09091 Arabidops
102	30	30.9	39	7	ADI23260	Novel hum	175	29	29.9	44	5	ABG09090	Abg09090 Human col
103	30	30.9	39	8	ADH74262	Human sec	176	29	29.9	44	5	ABU04320	Abu04320 Human col
104	30	30.9	39	8	ADH74262	Human sec	177	29	29.9	44	6	ABU61907	Abu61907 Mouse gly
105	30	30.9	39	8	ADM57320	Anti-IR f	178	29	29.9	45	3	ABG59379	Agag59379 Arabidops
106	30	30.9	40	2	AAW74791	Human sec	179	29	29.9	46	4	ABO50392	Abos0392 Human sec
107	30	30.9	40	5	ABG95241	Human nov	180	29	29.9	46	6	ABO44649	Abos44649 Novel hum
108	30	30.9	40	5	ABO34435	Region of	181	29	29.9	46	7	ABO26129	Abos26129 Human pro
109	30	30.9	40	7	ADH32063	alphaA-in	182	29	29.9	47	3	ABG44907	Abg44907 Human sec
110	30	30.9	40	7	ADI23096	Human sec	183	29	29.9	47	4	ABG02562	Abg02562 Novel hum
111	30	30.9	40	8	ADH74098	Human imm	184	29	29.9	47	5	AAW47156	Aaw47156 Modular e
112	30	30.9	41	4	AAW85472	Human mus	185	29	29.9	47	5	AAW47171	Aaw47171 Modular e
113	30	30.9	42	5	ABP29414	Streptoco	186	29	29.9	47	8	ABO54785	Abos4785 Human gen
114	30	30.9	46	6	ABU12775	Novel hum	187	29	29.9	48	3	ABO54785	Abos4785 Human gen
115	30	30.9	46	6	ABU12775	Novel hum	188	29	29.9	48	3	ABO54785	Abos4785 Human gen
116	30	30.9	46	8	ADJ28801	Human mus	189	29	29.9	48	3	ABO54785	Abos4785 Human gen
117	30	30.9	46	8	ADJ28801	Human mus	190	29	29.9	48	3	ABO54785	Abos4785 Human gen
118	29.5	30.4	32	5	ABP33068	Human ORF	191	29	29.9	48	4	ABG73655	Abg73655 Lung canc
119	29.5	30.4	33	5	ABP30850	Streptoco	192	29	29.9	49	3	ABG73655	Abg73655 Lung canc
120	29.5	30.4	33	5	ABP28726	Streptoco	193	29	29.9	50	3	ABG73655	Abg73655 Lung canc
121	29.5	30.4	44	3	AAW60246	Arabidops	194	29	29.9	50	3	ABG73655	Abg73655 Lung canc
122	29.5	30.4	48	4	AAO11779	Arabidops	195	29	29.9	50	3	ABG73655	Abg73655 Lung canc
123	29	29.9	20	7	ADW42197	Human pol	196	29	29.9	50	5	ABP29303	Abp29303 Streptoco
124	29	29.9	20	7	ADW42197	Human pol	197	29	29.9	50	5	ABP29303	Abp29303 Streptoco
125	29	29.9	21	1	ADP75643	T-cell ep	198	29	29.9	50	7	ADG90406	Adg90406 Novel hum
126	29	29.9	21	1	ADP75643	T-cell ep	199	29	29.9	50	7	ADG90406	Adg90406 Novel hum
127	29	29.9	26	8	ADI40693	Immunomod	200	28.5	29.4	30	4	AAW73727	Aaw73727 Peptide #
128	29	29.9	26	8	ADI40693	Immunomod	201	28.5	29.4	30	4	AAW73727	Aaw73727 Peptide #
129	29	29.9	28	2	AAW72756	Human PDZ	202	28.5	29.4	30	5	ABG5472	Abg5472 Human liv
130	29	29.9	28	4	ABR41309	Peptide #	203	28	28.9	42	2	AAW78190	Aaw78190 Amino aci
131	29	29.9	28	4	ABR41309	Peptide #	204	28	28.9	19	2	AAW78190	Aaw78190 Amino aci
132	29	29.9	28	4	ABR41309	Peptide #	205	28	28.9	19	4	AAW78190	Aaw78190 Amino aci
133	29	29.9	28	4	ABR41309	Peptide #	206	28	28.9	19	4	AAW78190	Aaw78190 Amino aci
134	29	29.9	28	4	ABR41309	Peptide #	207	28	28.9	19	4	AAW78190	Aaw78190 Amino aci
135	29	29.9	28	5	ABG44713	Human bra	208	28	28.9	19	4	AAW78190	Aaw78190 Amino aci
136	29	29.9	28	5	ABG44713	Human bra	209	28	28.9	19	4	AAW78190	Aaw78190 Amino aci
137	29	29.9	28	5	ABG44713	Human bra	210	28	28.9	19	4	AAW78190	Aaw78190 Amino aci
138	29	29.9	29	3	ABP27777	Sequence	211	28	28.9	23	2	AAW78190	Aaw78190 Amino aci
139	29	29.9	31	2	AAW19976	Fibronect	212	28	28.9	23	2	AAW78190	Aaw78190 Amino aci
140	29	29.9	31	2	AAW19976	Fibronect	213	28	28.9	23	2	AAW78190	Aaw78190 Amino aci
141	29	29.9	31	3	ABP39245	Human fib	214	28	28.9	23	4	ABW15118	Abw15118 Peptide #
142	29	29.9	32	8	ABO55532	Human gen	215	28	28.9	23	4	ABW15118	Abw15118 Peptide #
143	29	29.9	33	4	AAW42284	Human bre	216	28	28.9	23	4	ABW15118	Abw15118 Peptide #
144	29	29.9	33	4	AAW42284	Human bre	217	28	28.9	23	4	ABW15118	Abw15118 Peptide #
145	29	29.9	33	5	ABG62243	Human rep	218	28	28.9	23	4	ABW15118	Abw15118 Peptide #
146	29	29.9	33	5	ABG62243	Human rep	219	28	28.9	23	4	ABW15118	Abw15118 Peptide #
147	29	29.9	33	8	ADI40691	Hybrid po	220	28	28.9	23	4	ABW15118	Abw15118 Peptide #
148	29	29.9	34	2	AAW30694	Nef/SH3 d	221	28	28.9	23	4	ABW15118	Abw15118 Peptide #
149	29	29.9	34	3	AAW30694	Nef/SH3 d	222	28	28.9	23	4	ABW15118	Abw15118 Peptide #
150	29	29.9	34	3	AAW30694	Nef/SH3 d	223	28	28.9	23	4	ABW15118	Abw15118 Peptide #
151	29	29.9	34	3	AAW30694	Nef/SH3 d	224	28	28.9	23	4	ABW15118	Abw15118 Peptide #
152	29	29.9	34	3	AAW30694	Nef/SH3 d	225	28	28.9	23	4	ABW15118	Abw15118 Peptide #
153	29	29.9	34	3	AAW30694	Nef/SH3 d	226	28	28.9	23	4	ABW15118	Abw15118 Peptide #
154	29	29.9	34	3	AAW30694	Nef/SH3 d	227	28	28.9	23	4	ABW15118	Abw15118 Peptide #
155	29	29.9	34	3	AAW30694	Nef/SH3 d	228	28	28.9	23	4	ABW15118	Abw15118 Peptide #
156	29	29.9	34	3	AAW30694	Nef/SH3 d	229	28	28.9	23	4	ABW15118	Abw15118 Peptide #
157	29	29.9	34	3	AAW30694	Nef/SH3 d	230	28	28.9	23	4	ABW15118	Abw15118 Peptide #
158	29	29.9	34	3	AAW30694	Nef/SH3 d	231	28	28.9	23	4	ABW15118	Abw15118 Peptide #
159	29	29.9	34	3	AAW30694	Nef/SH3 d	232	28	28.9	23	4	ABW15118	Abw15118 Peptide #
160	29	29.9	34	3	AAW30694	Nef/SH3 d	233	28	28.9	23	4	ABW15118	Abw15118 Peptide #
161	29	29.9	34	3	AAW30694	Nef/SH3 d	234	28	28.9	23	4	ABW15118	Abw15118 Peptide #
162	29	29.9	34	3	AAW30694	Nef/SH3 d	235	28	28.9	23	4	ABW15118	Abw15118 Peptide #
163	29	29.9	34	3	AAW30694	Nef/SH3 d	236	28	28.9	23	4	ABW15118	Abw15118 Peptide #
164	29	29.9	34	3	AAW30694	Nef/SH3 d	237	28	28.9	23	4	ABW15118	Abw15118 Peptide #
165	29	29.9	34	3	AAW30694	Nef/SH3 d	238	28	28.9	23	4	ABW15118	Abw15118 Peptide #
166	29	29.9	34	3	AAW30694	Nef/SH3 d	239	28	28.9	23	4	ABW15118	Abw15118 Peptide #
167	29	29.9	34	3	AAW30694	Nef/SH3 d	240	28	28.9	23	4	ABW15118	Abw15118 Peptide #
168	29	29.9	34	3	AAW30694	Nef/SH3 d	241	28	28.9	23	4	ABW15118	Abw15118 Peptide #
169	29	29.9	34	3	AAW30694	Nef/SH3 d	242	28	28.9	23	4	ABW15118	Abw15118 Peptide #
170	29	29.9	34	3	AAW30694	Nef/SH3 d	243	28	28.9	23	4	ABW15118	Abw15118 Peptide #
171	29	29.9	34	3	AAW30694	Nef/SH3 d	244	28	28.9	23	4	ABW15118	Abw15118 Peptide #

245	28	28.9	38	2	AA13001	AA13001 Human sec
246	28	28.9	38	4	AA15186	Peptide #
247	28	28.9	38	4	ABB34179	Peptide #
248	28	28.9	38	4	AA27645	Peptide #
249	28	28.9	38	4	ABB29012	Peptide #
250	28	28.9	38	4	ABB19620	Protein #
251	28	28.9	38	4	AA67353	Human bon
252	28	28.9	38	4	AA54970	Human bra
253	28	28.9	38	4	ABG49016	Human liv
254	28	28.9	38	4	AA02925	Peptide #
255	28	28.9	38	5	ABG36988	Human pep
256	28	28.9	39	4	AA82762	Human imm
257	28	28.9	39	5	AA88731	Insulin/i
258	28	28.9	39	5	AA88749	Insulin/i
259	28	28.9	39	6	ADA03589	IGF-IR re
260	28	28.9	39	6	ADA03572	Insulin r
261	28	28.9	39	7	ADH94802	Insulin g
262	28	28.9	39	7	ADH94785	Insulin r
263	28	28.9	39	8	ADL67476	IGF-1R/IR
264	28	28.9	39	8	ADL67493	IGF-1R/IR
265	28	28.9	39	8	ADM37338	Anti-IGF-
266	28	28.9	39	8	ADM37321	Anti-IR f
267	28	28.9	40	6	AAE36707	Rat Ti-VA
268	28	28.9	40	7	ABW01763	Rat Ti-VA
269	28	28.9	40	8	ADM97081	Botulinum
270	28	28.9	41	4	AAU22274	Human car
271	28	28.9	41	4	AAE46242	Human car
272	28	28.9	42	2	AAW88776	Polypepti
273	28	28.9	42	4	ABB50609	Human sec
274	28	28.9	42	6	ABO44866	Novel hum
275	28	28.9	42	7	ABO26346	Protein a
276	28	28.9	43	7	ADF70043	Acna-type
277	28	28.9	44	4	ABB68452	Drosophil
278	28	28.9	45	3	AA32980	Arabidops
279	28	28.9	45	4	AA92489	Human dig
280	28	28.9	45	4	AA86098	Human imm
281	28	28.9	45	4	AAU22534	Novel hum
282	28	28.9	45	7	ADB32374	Human nov
283	28	28.9	47	2	AAW78184	Human sec
284	28	28.9	47	4	AAW13761	Peptide #
285	28	28.9	47	4	AA374606	Human col
286	28	28.9	47	4	ABB32694	Peptide #
287	28	28.9	47	4	AA26159	Peptide #
288	28	28.9	47	4	AA83381	Human imm
289	28	28.9	47	4	ABB27535	Human pep
290	28	28.9	47	4	ABB18183	Protein #
291	28	28.9	47	4	AA65894	Human bon
292	28	28.9	47	4	AA53516	Human bra
293	28	28.9	47	4	ABG47549	Human liv
294	28	28.9	47	4	AA01506	Peptide #
295	28	28.9	47	5	ABG35529	Human pep
296	28	28.9	48	4	AA851100	Human imm
297	28	28.9	48	5	ABP28234	Streptoco
298	28	28.9	48	5	ADD44478	Polypepti
299	28	28.9	50	3	AAQ08758	Arabidops
300	28	28.9	50	5	ADH32512	Yeast smo
301	28	28.9	50	6	ABM71782	Staphyloc
302	27.5	28.4	30	2	AA23781	N terminu
303	27.5	28.4	41	4	ABB16647	Human ner
304	27	27.8	19	2	AA50599	Resin bou
305	27	27.8	19	7	ADF14612	Rheumatoi
306	27	27.8	20	2	AAW42169	T-cell ep
307	27	27.8	21	2	AAW59015	Lactococc
308	27	27.8	21	2	AAW94873	N-termina
309	27	27.8	21	5	AAE20718	Human Mls
310	27	27.8	21	5	AAE21019	Human Icr
311	27	27.8	21	8	ADH89723	Cell pene
312	27	27.8	22	2	AA04241	Human par
313	27	27.8	22	7	ADI24804	Parathyro
314	27	27.8	23	2	AA04242	Human par
315	27	27.8	23	5	AA71457	Cobra C3
316	27	27.8	23	5	AA71459	Murine C3
317	27	27.8	23	7	ADI24805	Parathyro

318	27	27.8	23	8	ADQ35122
319	27	27.8	24	2	AAW04243
320	27	27.8	24	7	ADI24806
321	27	27.8	25	4	AAW04244
322	27	27.8	25	4	ABG09991
323	27	27.8	25	7	ADI24807
324	27	27.8	25	8	ABW79621
325	27	27.8	25	8	ADJ36098
326	27	27.8	26	2	AAW61646
327	27	27.8	26	2	AAW64665
328	27	27.8	26	2	AAW04245
329	27	27.8	26	3	AA12831
330	27	27.8	26	7	ADI24808
331	27	27.8	26	8	ADQ14464
332	27	27.8	27	2	AAW59014
333	27	27.8	27	2	AAW06348
334	27	27.8	27	2	AAW94872
335	27	27.8	27	2	AAW04227
336	27	27.8	27	2	AAW04246
337	27	27.8	27	2	AAW27328
338	27	27.8	27	4	AAW14225
339	27	27.8	27	4	ABB33172
340	27	27.8	27	4	AAW26635
341	27	27.8	27	4	ABB28000
342	27	27.8	27	4	ABB18637
343	27	27.8	27	4	AAW66356
344	27	27.8	27	4	AAW53968
345	27	27.8	27	4	ABG48022
346	27	27.8	27	4	AAW01957
347	27	27.8	27	5	ABG36004
348	27	27.8	27	7	ADI24809
349	27	27.8	27	7	ADI24803
350	27	27.8	28	2	AAW04226
351	27	27.8	28	2	AAW04248
352	27	27.8	28	2	AAW04247
353	27	27.8	28	2	AAW04240
354	27	27.8	28	2	AAW50589
355	27	27.8	28	2	AAW50592
356	27	27.8	28	7	ADI24810
357	27	27.8	28	7	ADI24802
358	27	27.8	28	7	ADI24827
359	27	27.8	28	7	ADI24813
360	27	27.8	29	2	AAW04225
361	27	27.8	29	2	AAW04228
362	27	27.8	29	7	ADI24801
363	27	27.8	29	7	ADI24811
364	27	27.8	30	2	AAW04224
365	27	27.8	30	2	AAW04229
366	27	27.8	30	2	AAW50601
367	27	27.8	30	4	ABB38725
368	27	27.8	30	4	AAW32194
369	27	27.8	30	4	AAW71913
370	27	27.8	30	4	AAW59358
371	27	27.8	30	4	ABG53596
372	27	27.8	30	5	ABG41727
373	27	27.8	30	5	AAU84870
374	27	27.8	30	7	ADI24812
375	27	27.8	30	7	ADI24800
376	27	27.8	31	2	AAW04179
377	27	27.8	31	2	AAW04209
378	27	27.8	31	2	AAW04254
379	27	27.8	31	2	AAW04192
380	27	27.8	31	2	AAW04208
381	27	27.8	31	2	AAW04238
382	27	27.8	31	2	AAW04255
383	27	27.8	31	2	AAW04186
384	27	27.8	31	2	AAW04187
385	27	27.8	31	2	AAW04250
386	27	27.8	31	2	AAW04178
387	27	27.8	31	2	AAW04184
388	27	27.8	31	2	AAW04199
389	27	27.8	31	2	AAW04202
390	27	27.8	31	2	AAW04210

Adq35122	Novel pep
Aay04243	Human par
Adi24806	Parathyro
Aay04244	Human par
ABG09991	Novel hum
ADI24807	Parathyro
ABM79621	M tubercu
ADJ36098	Self-coal
AAW61646	v-Src ATP
AAW64665	Synthetic
AAW04245	Human par
AA12831	V-Src ATP
ADI24808	Parathyro
ADQ14464	C2 H2 typ
AAW59014	Lactococc
AAW06348	Mycelloph
AAW94872	N-termina
AAW04227	Human par
AAW04246	Human par
AAW27328	Human C9
AAW14225	Peptide #
ABB33172	Peptide #
ABB26635	Peptide #
ABB28000	Human pep
ABB18637	Protein #
AAW66356	Human bon
AAW53968	Human bra
ABG48022	Human liv
AAW01957	Peptide #
ABG36004	Human pep
ADI24809	Parathyro
ADI24803	Parathyro
Aay04226	Human par
Aay04248	Human par
Aay04240	Human par
Aay50589	Resin bou
Aay50592	Resin bou
Aay50592	Resin bou
ADI24810	Parathyro
ADI24802	Parathyro
ADI24827	Parathyro
ADI24813	Parathyro
Aay04225	Human par
Aay04228	Human par
Aay04229	Human par
ADI24801	Parathyro
ADI24811	Parathyro
Aay04224	Human par
Aay04229	Human par
Aay50601	Resin bou
ABB38725	Peptide #
AAW32194	Peptide #
AAW71913	Human bon
AAW59358	Human bra
ABG53596	Human liv
ABG41727	Human pep
AAU84870	Human MAR
ADI24812	Parathyro
AAW04179	Human par
AAW04179	Human par
AAW04209	Human par
AAW04254	Human par
AAW04192	Human par
AAW04208	Human par
AAW04238	Human par
AAW04255	Human par
AAW04186	Human par
AAW04187	Human par
AAW04250	Human par
AAW04178	Human par
AAW04184	Human par
AAW04199	Human par
AAW04202	Human par
AAW04210	Human par

391	27	27.8	31	2	AA04256	Human par	Aay04256	Human par	464	27	27.8	31	7	ADI24785	Parathyro
392	27	27.8	31	2	AA04258	Human par	Aay04258	Human par	465	27	27.8	31	7	ADI24789	Parathyro
393	27	27.8	31	2	AA04185	Human par	Aay04185	Human par	466	27	27.8	31	7	ADI24754	Parathyro
394	27	27.8	31	2	AA04198	Human par	Aay04198	Human par	467	27	27.8	31	7	ADI24822	Parathyro
395	27	27.8	31	2	AA04213	Human par	Aay04213	Human par	468	27	27.8	31	7	ADI24833	Parathyro
396	27	27.8	31	2	AA04220	Human par	Aay04220	Human par	469	27	27.8	31	7	ADI24836	Parathyro
397	27	27.8	31	2	AA04183	Human par	Aay04183	Human par	470	27	27.8	31	7	ADI24788	Parathyro
398	27	27.8	31	2	AA04211	Human par	Aay04211	Human par	471	27	27.8	32	4	AA62184	Human gen
399	27	27.8	31	2	AA04259	Human par	Aay04259	Human par	472	27	27.8	32	5	ABG6383	Human alb
400	27	27.8	31	2	AA04195	Human par	Aay04195	Human par	473	27	27.8	32	8	ADL76848	Human f
401	27	27.8	31	2	AA04261	Human par	Aay04261	Human par	474	27	27.8	33	4	AA677644	Human col
402	27	27.8	31	2	AA04176	Human par	Aay04176	Human par	475	27	27.8	34	1	AA92220	Peptide s
403	27	27.8	31	2	AA04182	Human par	Aay04182	Human par	476	27	27.8	34	2	AA04218	Human par
404	27	27.8	31	2	AA04194	Human par	Aay04194	Human par	477	27	27.8	34	2	AA50585	Resin bou
405	27	27.8	31	2	AA04222	Human par	Aay04222	Human par	478	27	27.8	34	3	AA53927	Human col
406	27	27.8	31	2	AA04180	Human par	Aay04180	Human par	479	27	27.8	34	4	AAU18637	Human lun
407	27	27.8	31	2	AA04193	Human par	Aay04193	Human par	480	27	27.8	34	7	ADB33261	Human nov
408	27	27.8	31	2	AA04203	Human par	Aay04203	Human par	481	27	27.8	34	7	ADI24794	Parathyro
409	27	27.8	31	2	AA04257	Human par	Aay04257	Human par	482	27	27.8	35	3	AA589302	Core poly
410	27	27.8	31	2	AA04197	Human par	Aay04197	Human par	483	27	27.8	35	4	AA577704	Core poly
411	27	27.8	31	2	AA04200	Human par	Aay04200	Human par	484	27	27.8	35	4	AB500710	Core poly
412	27	27.8	31	2	AA04177	Human par	Aay04177	Human par	485	27	27.8	35	4	ABB02187	Virial cor
413	27	27.8	31	2	AA04181	Human par	Aay04181	Human par	486	27	27.8	35	4	AAU13257	Human nov
414	27	27.8	31	2	AA04196	Human par	Aay04196	Human par	487	27	27.8	35	4	AAU20775	Human nov
415	27	27.8	31	2	AA04201	Human par	Aay04201	Human par	488	27	27.8	35	5	ADH32768	Yeast smc
416	27	27.8	31	2	AA04212	Human par	Aay04212	Human par	489	27	27.8	36	2	AA558185	[lys18]-h
417	27	27.8	31	2	AA050575	Resin bou	Aay50575	Resin bou	490	27	27.8	36	4	AAE03915	Human gen
418	27	27.8	31	2	AA050571	Resin bou	Aay50571	Resin bou	491	27	27.8	36	4	ABB16886	Human ner
419	27	27.8	31	2	AA050579	Resin bou	Aay50579	Resin bou	492	27	27.8	36	8	ABO55022	Human gen
420	27	27.8	31	2	AA050570	Resin bou	Aay50570	Resin bou	493	27	27.8	37	3	AA16673	Bacteriop
421	27	27.8	31	2	AA050584	Resin bou	Aay50584	Resin bou	494	27	27.8	37	3	AA45168	Human sec
422	27	27.8	31	2	AA050569	Resin bou	Aay50569	Resin bou	495	27	27.8	37	4	AA664430	Human sec
423	27	27.8	31	2	AA050574	Resin bou	Aay50574	Resin bou	496	27	27.8	37	8	ADF45218	Human INP
424	27	27.8	31	2	AA050577	Resin bou	Aay50577	Resin bou	497	27	27.8	38	3	AA10619	Human SAP
425	27	27.8	31	2	AA050576	Resin bou	Aay50576	Resin bou	498	27	27.8	38	7	ABO23537	Borrelia
426	27	27.8	31	2	AA050578	Resin bou	Aay50578	Resin bou	499	27	27.8	38	7	ADF76747	Novel hum
427	27	27.8	31	2	AA050573	Resin bou	Aay50573	Resin bou	500	27	27.8	38	8	ADM37301	Anti-IR f
428	27	27.8	31	2	AA050568	Resin bou	Aay50568	Resin bou							
429	27	27.8	31	2	AA050572	Resin bou	Aay50572	Resin bou							
430	27	27.8	31	7	ADI24768	Parathyro	Adi24768	Parathyro							
431	27	27.8	31	7	ADI24763	Parathyro	Adi24763	Parathyro							
432	27	27.8	31	7	ADI24751	Parathyro	Adi24751	Parathyro							
433	27	27.8	31	7	ADI24757	Parathyro	Adi24757	Parathyro							
434	27	27.8	31	7	ADI24761	Parathyro	Adi24761	Parathyro							
435	27	27.8	31	7	ADI24772	Parathyro	Adi24772	Parathyro							
436	27	27.8	31	7	ADI24787	Parathyro	Adi24787	Parathyro							
437	27	27.8	31	7	ADI24831	Parathyro	Adi24831	Parathyro							
438	27	27.8	31	7	ADI24774	Parathyro	Adi24774	Parathyro							
439	27	27.8	31	7	ADI24758	Parathyro	Adi24758	Parathyro							
440	27	27.8	31	7	ADI24770	Parathyro	Adi24770	Parathyro							
441	27	27.8	31	7	ADI24832	Parathyro	Adi24832	Parathyro							
442	27	27.8	31	7	ADI24834	Parathyro	Adi24834	Parathyro							
443	27	27.8	31	7	ADI24753	Parathyro	Adi24753	Parathyro							
444	27	27.8	31	7	ADI24759	Parathyro	Adi24759	Parathyro							
445	27	27.8	31	7	ADI24775	Parathyro	Adi24775	Parathyro							
446	27	27.8	31	7	ADI24752	Parathyro	Adi24752	Parathyro							
447	27	27.8	31	7	ADI24760	Parathyro	Adi24760	Parathyro							
448	27	27.8	31	7	ADI24776	Parathyro	Adi24776	Parathyro							
449	27	27.8	31	7	ADI24762	Parathyro	Adi24762	Parathyro							
450	27	27.8	31	7	ADI24778	Parathyro	Adi24778	Parathyro							
451	27	27.8	31	7	ADI24829	Parathyro	Adi24829	Parathyro							
452	27	27.8	31	7	ADI24773	Parathyro	Adi24773	Parathyro							
453	27	27.8	31	7	ADI24798	Parathyro	Adi24798	Parathyro							
454	27	27.8	31	7	ADI24756	Parathyro	Adi24756	Parathyro							
455	27	27.8	31	7	ADI24779	Parathyro	Adi24779	Parathyro							
456	27	27.8	31	7	ADI24830	Parathyro	Adi24830	Parathyro							
457	27	27.8	31	7	ADI24755	Parathyro	Adi24755	Parathyro							
458	27	27.8	31	7	ADI24771	Parathyro	Adi24771	Parathyro							
459	27	27.8	31	7	ADI24786	Parathyro	Adi24786	Parathyro							
460	27	27.8	31	7	ADI24835	Parathyro	Adi24835	Parathyro							
461	27	27.8	31	7	ADI24769	Parathyro	Adi24769	Parathyro							
462	27	27.8	31	7	ADI24777	Parathyro	Adi24777	Parathyro							
463	27	27.8	31	7	ADI24784	Parathyro	Adi24784	Parathyro							

## ALIGNMENTS

## RESULT 1

AAR43461

ID AAR43461 standard; peptide; 22 AA.

XX AAR43461;

XX

DT 25-MAR-2003 (revised)

DT 12-MAY-1994 (first entry)

XX

DE Ro/SSA epitope 257.

XX

KW Linear; epitope; 60 kD; Ro/SSA; La/SSB; autoantigen; E/F; G; 70 kD;

KW nuclear ribonucleoprotein; rRNP; Sm B/B'; polypeptide; antigen; D;

KW systemic lupus erythematosus; SLE; autoantibody; U4/U6; U5; B; B';

KW RNA polymerase III; U1; U2; Sjogrens syndrome; SS; human; vaccine; ss.

XX

OS Homo sapiens.

XX

FN WO9321223-A1.

XX

PD 28-OCT-1993.

XX

PF 13-APR-1993; 93WO-US003484.

XX

PR 13-APR-1992; 92US-00867819.

XX

PA (OKLA ) UNIV OKLAHOMA STATE.

XX

PI Harley JB;

XX

DR WPI; 1993-351658/44.



XX New linear epitope(s) for human auto-antibodies - from the Ro/SSA, La/SSB  
 PT and Sm B/B' antigens and ribo:nucleoprotein, used for diagnosing and  
 PT treating auto-immune disorders e.g. systemic lupus erythematosus.  
 XX  
 PS Claim 1; Page 31; 43pp; English.  
 XX  
 CC The sequences given in AAR43391-562 are linear epitopes which are derived  
 CC from the 60 kD Ro/SSA peptide, the La/SSB autoantigen, the 70 kD nuclear  
 CC ribonucleoprotein (nRNP) and the Sm B/B' polypeptide. These antigens are  
 CC common in systemic lupus erythematosus (SLE) and closely related  
 CC disorders. The Ro/SSA family of proteins has been shown to have several  
 CC molecular forms which are defined by the molecular weight of the antigen  
 CC identified. The major form has a molecular weight of 60 kD and two  
 CC additional forms have molecular weights of 52 and 54 kD. La/SSB is also a  
 CC member of this group of autoantibodies and binds small RNAs with a  
 CC polypyridine terminus. La/SSB is bound by a third of the anti-Ro/SSA  
 CC precipitin positive sera. La/SSB has been shown to be a 46-50 kD  
 CC monomeric phosphoprotein which associates with RNA polymerase III  
 CC transcripts. Anti-Sm antibodies precipitate snRNPs containing the U1, U2,  
 CC U4/U6 and U5 RNA. Anti-Sm antibodies may be directed against one or a  
 CC combination of the polypeptides: B (26 kD), B' (27 kD), D (13 kD), E/F  
 CC (11 kD doublet) and G (less than 10 kD). These epitopes may be used for  
 CC preventing, treating or screening autoimmune disorders, especially SLE or  
 CC Sjogrens syndrome (SS). They bind to a human autoantibody and may  
 CC therefore be used as vaccines. (Updated on 25-MAR-2003 to correct PN  
 CC field.)  
 XX  
 CC Sequence 22 AA;

Query Match 39.2%; Score 38; DB 2; Length 22;  
 Best Local Similarity 53.3%; Pred. No. 21; Mismatches 0; Gaps 0;  
 Matches 8; Conservative 2; Indels 5;

QY 3 NHLNSKIAFKIVSQE 17  
 ||||| : : : : :  
 Db 5 NHLKSEVWKALQE 19

RESULT 2  
 AAB27169  
 ID AAB27169 standard; protein; 46 AA.

XX AAB27169;

XX 27-FEB-2001 (first entry)

XX Sendai virus partial protein sequence SEQ ID NO: 17.

XX Negative stranded RNA virus; vaccine; attenuated virus; RSV; PIV;  
 XX measles; respiratory syncytial virus; parainfluenza virus.

XX Sendai virus.

XX WO200061737-A2.

XX 19-OCT-2000.

XX 12-APR-2000; 2000WO-US009695.

XX 13-APR-1999; 99US-0129006P.

XX (USSH ) US DEPT HEALTH & HUMAN SERVICES.

XX Murphy BR, Collins PL, Durbin AP, Skiadopoulos MH;

XX WPI; 2000-687044/67.

XX Producing attenuated negative stranded RNA virus vaccines from cloned  
 PT sequences, useful for immunizing against e.g. respiratory syncytial  
 PT virus, human parainfluenza virus, Sendai virus Newcastle disease virus,  
 PT mumps virus and measles virus.

PS Example 1; Page 62; 137pp; English.

XX The present invention is concerned with producing vaccines against  
 CC negative stranded RNA viruses. These viruses include measles, respiratory  
 CC syncytial virus (RSV) and parainfluenza virus (PIV) in particular. The  
 CC method of the invention comprises the production of a mutated form of the  
 CC virus which attenuates the strain and enables it to be used as a vaccine.  
 CC The present sequence comprises a partial viral protein sequence

XX Sequence 46 AA;

Query Match 36.1%; Score 35; DB 3; Length 46;  
 Best Local Similarity 28.6%; Pred. No. 1.7e+02;  
 Matches 4; Conservative 8; Mismatches 2; Indels 0; Gaps 0;

QY 5 LNSKIAFKIVSQEP 18  
 ||: : : : :  
 Db 1 LDKQVLYRVNQP 14

RESULT 3  
 AABM72186  
 ID AABM72186 standard; protein; 45 AA.

XX AABM72186;

XX 20-NOV-2003 (first entry)

XX Staphylococcus aureus protein #1426.

XX Antibacterial; vaccine; gene therapy; infection; sepsis; diagnosis;  
 XX enzymatic assay; antibiotic target.

XX Staphylococcus aureus.

XX WO200294868-A2.

XX 28-NOV-2002.

XX 27-MAR-2002; 2002WO-IB002637.

XX 27-MAR-2001; 2001GB-00007661.

XX (CHIR-) CHIRON SPA.

XX Massignani V, Mora M, Scarselli M;

XX WPI; 2003-120786/11.

XX N-PSDB; ACF73746.

XX New Staphylococcus aureus protein, useful as a vaccine for treating or  
 PT preventing Staphylococcal infection, specifically an infection caused by  
 S. aureus, e.g. sepsis.

XX Claim 1; SEQ ID NO 2852; 49pp; English.

XX The invention relates to novel genes and encoded proteins from  
 CC Staphylococcus aureus. A composition comprising the S. aureus protein, a  
 CC nucleic acid encoding the protein, or an antibody to the protein, is  
 CC useful as a pharmaceutical, particularly as a vaccine for treating or  
 CC preventing infection due to Staphylococcus bacteria, specifically an  
 CC infection caused by S. aureus. The composition is particularly useful for  
 CC treating or preventing sepsis in a patient. The composition can also be  
 CC used for diagnostics. The protein is also used in an assay for enzymatic  
 CC studies and as a target for antibiotics. This sequence represents one of  
 CC the novel S. aureus proteins of the invention

XX Sequence 45 AA;

Query Match 35.1%; Score 34; DB 6; Length 45;  
 Best Local Similarity 62.5%; Pred. No. 2.6e+02;  
 Matches 5; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

Qy 2 PNHLNSKI 9  
 Db 16 PNHLNTDV 23

## RESULT 4

AAAB27167  
 ID AAB27167 standard; protein; 46 AA.

AC AAB27167;

DT 27-FEB-2001 (first entry)

DE HPIV3 partial protein sequence SEQ ID NO: 15.

KW Negative stranded RNA virus; vaccine; attenuated virus; RSV; PIV;  
 KW measles; respiratory syncytial virus; parainfluenza virus.

OS Human parainfluenza virus.

PN WO200061737-A2.

PD 19-OCT-2000.

PF 12-APR-2000; 2000WO-US009695.

PR 13-APR-1999; 99US-0129006P.

PA (USSH ) US DEPT HEALTH & HUMAN SERVICES.

PI Murphy BR, Collins PL, Durbin AP, Skiadopoulos MH;

DR WPI; 2000-687044/67.

PT Producing attenuated negative stranded RNA virus vaccines from cloned  
 PT sequences, useful for immunizing against e.g. respiratory syncytial  
 PT virus, human parainfluenza virus, Sendai virus Newcastle disease virus,  
 PT mumps virus and measles virus.

PS Example 1; Page 62; 137pp; English.

CC The present invention is concerned with producing vaccines against  
 CC negative stranded RNA viruses. These viruses include measles, respiratory  
 CC syncytial virus (RSV) and parainfluenza virus (PIV) in particular. The  
 CC method of the invention comprises the production of a mutated form of the  
 CC virus which attenuates the strain and enables it to be used as a vaccine.  
 CC The present sequence comprises a partial viral protein sequence

SQ Sequence 46 AA;

Query Match 35.1%; Score 34; DB 3; Length 46;

Best Local Similarity 35.7%; Pred. No. 2.6e+02;

Matches 5; Conservative 6; Mismatches 3; Indels 0; Gaps 0;

Qy 5 LNSKIAFKIVSQEP 18

Db 1 LDRSVLYRMNQEP 14

## RESULT 5

AAU21220

ID AAU21220 standard; protein; 49 AA.

AC AAU21220;

DT 17-DEC-2001 (first entry)

DE Human novel foetal antigen, SEQ ID NO 1464.

KW Human; foetal tissue antigen; antiinflammatory; neuroprotective;  
 KW immunomodulator; cardiovascular; cytostatic; nephrothropic;  
 KW cardiovascular; autoimmune disease; rheumatoid arthritis;  
 KW hyperproliferative disorder; breast neoplasm; cancer;

KW cardiovascular disorder; cardiac arrest; cerebrovascular disorder;  
 KW cerebral ischaemia; angiogenesis; nervous system disorder;  
 KW Alzheimer's disease; infection; ocular disorder; corneal infection;  
 KW wound healing; epithelial cell proliferation; food additive.

OS Homo sapiens.

PN WO200155312-A2.

XX 02-AUG-2001.

PF 17-JAN-2001; 2001WO-US001321.

PR 31-JAN-2000; 2000US-0179065P.

PR 04-FEB-2000; 2000US-0180628P.

PR 24-FEB-2000; 2000US-0184664P.

PR 02-MAR-2000; 2000US-0186350P.

PR 16-MAR-2000; 2000US-0189874P.

PR 17-MAR-2000; 2000US-0190076P.

PR 18-APR-2000; 2000US-0198123P.

PR 19-MAY-2000; 2000US-0205515P.

PR 07-JUN-2000; 2000US-0209467P.

PR 28-JUN-2000; 2000US-0214886P.

PR 30-JUN-2000; 2000US-0215135P.

PR 07-JUL-2000; 2000US-0216647P.

PR 07-JUL-2000; 2000US-0216880P.

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PR 26-JUL-2000; 2000US-0220964P.

PR 14-AUG-2000; 2000US-0224518P.

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PR 14-AUG-2000; 2000US-0225757P.

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PR 18-AUG-2000; 2000US-0225759P.

PR 22-AUG-2000; 2000US-0226279P.

PR 22-AUG-2000; 2000US-0226681P.

PR 22-AUG-2000; 2000US-0226868P.

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PR 01-SEP-2000; 2000US-0229344P.

PR 01-SEP-2000; 2000US-0229345P.

PR 05-SEP-2000; 2000US-0229509P.

PR 06-SEP-2000; 2000US-0229513P.

PR 06-SEP-2000; 2000US-0230437P.

PR 06-SEP-2000; 2000US-0230438P.

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PR 08-SEP-2000; 2000US-0231243P.

PR 08-SEP-2000; 2000US-0231244P.

PR 08-SEP-2000; 2000US-0231413P.

PR 08-SEP-2000; 2000US-0231414P.

PR 08-SEP-2000; 2000US-0232080P.

PR 08-SEP-2000; 2000US-0232081P.

PR 12-SEP-2000; 2000US-0231968P.

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PR 14-SEP-2000; 2000US-0232398P.

PR 14-SEP-2000; 2000US-0232399P.

PR 14-SEP-2000; 2000US-0232400P.

PR 14-SEP-2000; 2000US-0232401P.

PR 14-SEP-2000; 2000US-0233063P.

PR 14-SEP-2000; 2000US-0233064P.

PR 14-SEP-2000; 2000US-0233065P.

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PR 08-NOV-2000; 2000US-0246476P.
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PR 08-NOV-2000; 2000US-0246610P.
PR 08-NOV-2000; 2000US-0246611P.
PR 17-NOV-2000; 2000US-0246613P.
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PR 17-NOV-2000; 2000US-0249245P.
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PR 17-NOV-2000; 2000US-0249265P.
PR 17-NOV-2000; 2000US-0249297P.
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PR 05-DEC-2000; 2000US-0251988P.
PR 05-DEC-2000; 2000US-0256719P.
PR 06-DEC-2000; 2000US-0251479P.
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PR 21-SEP-2000; 2000US-0251990P.
PR 11-DEC-2000; 2000US-0254097P.
PR 05-JAN-2001; 2001US-0259678P.
XX (HUMA-) HUMAN GENOME SCI INC.
XX Rosen CA, Barash SC, Ruben SM;
XX WPI; 2001-488782/53.
XX N-PSDB; AAS34040.
XX New polynucleotides and polypeptides for diagnosing, treating, preventing
PT or prognosing e.g. diseases or disorders of the nervous, musculoskeletal,
PT excretory, gastrointestinal, reproductive, and respiratory systems.
XX Claim 11; SEQ ID NO 1464; 642pp; English.
XX The invention relates to novel nucleic acids encoding novel human foetal
CC antigens. The nucleic acids and proteins are used to prevent, treat (e.g.
CC by gene therapy) or ameliorate a medical condition in e.g. humans, mice,
CC rabbits, goats, horses, cats, dogs, chickens or sheep. They are also used
CC in diagnosing a pathological condition or susceptibility to a
CC pathological condition. The antibodies to the antigens and in diagnostic
CC in alleviating symptoms associated with the disorders and in diagnostic
CC immunoassays e.g. radioimmunoassays or enzyme linked immunosorbent assays
CC (ELISA). Disorders which are diagnosed or treated include autoimmune
CC diseases e.g. rheumatoid arthritis, hyperproliferative disorders e.g.
CC neoplasms of the breast or liver, cardiovascular disorders e.g. cardiac
CC arrest, cerebrovascular disorders e.g. cerebral ischaemia, angiodenesis,
CC nervous system disorders e.g. Alzheimer's disease, infections caused by
CC bacteria, viruses and fungi and ocular disorders e.g. corneal infection.
CC The polypeptides can also be used to aid wound healing and epithelial
CC cell proliferation, to prevent skin aging due to sunburn, to maintain
CC organs before transplantation, for supporting cell culture of primary
CC tissues, to regenerate tissues and in chemotaxis. The polypeptides can
CC also be used as a food additive or preservative to increase or decrease
CC storage capabilities, fat content, lipid, protein, carbohydrate,
CC vitamins, minerals, cofactors and other nutritional components. Numerous
CC examples of diseases and disorders treated by the nucleic acids and
CC proteins are given in the specification. The present sequence represents
CC a foetal antigen of the invention. Note: The sequence data for this
CC patent did not form part of the printed specification, but was obtained
Query Match 35.1%; Score 34; DB 4; Length 49;
Best Local Similarity 70.0%; Pred. No. 2.8e+02;
Matches 7; Conservative 0; Mismatches 3; Indels 0; Gaps 0;
Qy 2 PNHLSKIAF 11
Db 3 PFHNSKIKE 12
RESULT 6
AAM33005
ID AAM33005 standard; protein; 26 AA.
AC AAM33005;
XX 17-OCT-2001 (first entry)
DT 17-OCT-2001 (first entry)
DE Peptide #7042 encoded by probe for measuring placental gene expression.
XX Probe; microarray; human; placenta; antenatal diagnosis;
XX genetic disorder.
XX Homo sapiens.
XX WO200157272-A2.
XX 09-AUG-2001.
XX 30-JAN-2001; 2001WO-US000663.
XX
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PR 04-FEB-2000; 2000US-0180312P.
PR 26-MAY-2000; 2000US-0207456P.
PR 30-JUN-2000; 2000US-00608408.
PR 03-AUG-2000; 2000US-00632366.
PR 21-SEP-2000; 2000US-0234687P.
PR 27-SEP-2000; 2000US-0236359P.
PR 04-OCT-2000; 2000GB-00024263.
XX (MOLE-) MOLECULAR DYNAMICS INC.
XX Penn SG, Hanzel DK, Chen W, Rank DR;
XX WPI; 2001-488997/53.
XX Human genome-derived single exon nucleic acid probes useful for analyzing
XX gene expression in human placenta.
XX Claim 27; SEQ ID NO 33274; 654pp; English.
XX The present invention relates to single exon nucleic acid probes (SENP:
XX see AAI31315-AAI57546). The present sequence is a peptide encoded by one
XX such probe. The probes are useful for producing a microarray for
XX predicting, measuring and displaying gene expression in samples derived
XX from human placenta. The probes are useful for antenatal diagnosis of
XX human genetic disorders
XX Sequence 26 AA;
Query Match 34.0%; Score 33; DB 4; Length 26;
Best Local Similarity 43.8%; Pred. No. 2e+02;
Matches 7; Conservative 2; Mismatches 7; Indels 0; Gaps 0;
QY 3 NHIKSKIAPKIVSQEP 18
DB 2 NTLERKTPILQIQEP 17
RESULT 7
AAM72776
ID AAM72776 standard; protein; 26 AA.
XX AAM72776;
XX 06-NOV-2001 (first entry)
XX Human bone marrow expressed probe encoded protein SEQ ID NO: 33082.
XX Human; bone marrow expressed exon; gene expression analysis; probe;
XX microarray; cancer; leukaemia; lymphoma; myeloma.
XX Homo sapiens.
XX WO200157276-A2.
XX 09-AUG-2001.
XX 30-JAN-2001; 2001WO-US000668.
XX 04-FEB-2000; 2000US-0180312P.
XX 26-MAY-2000; 2000US-0207456P.
XX 30-JUN-2000; 2000US-00608408.
XX 03-AUG-2000; 2000US-00632366.
XX 21-SEP-2000; 2000US-0234687P.
XX 27-SEP-2000; 2000US-0236359P.
XX 04-OCT-2000; 2000GB-00024263.
XX (MOLE-) MOLECULAR DYNAMICS INC.
XX Penn SG, Hanzel DK, Chen W, Rank DR;
XX WPI; 2001-488990/53.
XX Human genome-derived single exon nucleic acid probes useful for analyzing

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PT gene expression in human bone marrow.
XX Example 4; SEQ ID NO 33082; 658pp + Sequence Listing; English.
XX The present invention provides a number of single exon nucleic acid
XX probes which are derived from genomic sequences expressed in the human
XX bone marrow. They can be used to measure gene expression in bone marrow
XX samples, which may enable the improved diagnosis and treatment of cancers
XX such as lymphoma, leukaemia and myeloma. The present sequence is a
XX protein encoded by one of the probes of the invention
XX Sequence 26 AA;
Query Match 34.0%; Score 33; DB 4; Length 26;
Best Local Similarity 43.8%; Pred. No. 2e+02;
Matches 7; Conservative 2; Mismatches 7; Indels 0; Gaps 0;
QY 3 NHIKSKIAPKIVSQEP 18
DB 2 NTLERKTPILQIQEP 17
RESULT 8
AAM60161
ID AAM60161 standard; protein; 26 AA.
XX AAM60161;
XX 05-NOV-2001 (first entry)
XX Human brain expressed single exon probe encoded protein SEQ ID NO: 32266.
XX Human; brain expressed exon; gene expression analysis; probe; microarray;
XX Alzheimer's disease; multiple sclerosis; schizophrenia; epilepsy; cancer.
XX Homo sapiens.
XX WO200157275-A2.
XX 09-AUG-2001.
XX 30-JAN-2001; 2001WO-US000667.
XX 04-FEB-2000; 2000US-0180312P.
XX 26-MAY-2000; 2000US-0207456P.
XX 30-JUN-2000; 2000US-00608408.
XX 03-AUG-2000; 2000US-00632366.
XX 21-SEP-2000; 2000US-0234687P.
XX 27-SEP-2000; 2000US-0236359P.
XX 04-OCT-2000; 2000GB-00024263.
XX (MOLE-) MOLECULAR DYNAMICS INC.
XX Penn SG, Hanzel DK, Chen W, Rank DR;
XX WPI; 2001-483446/52.
XX Single exon nucleic acid probes for analyzing gene expression in human
XX brains.
XX Example 4; SEQ ID NO 32266; 650pp + Sequence Listing; English.
XX The present invention provides a number of single exon nucleic acid
XX probes which are derived from genomic sequences expressed in the human
XX brain. They can be used to measure gene expression in brain cell samples,
XX which may enable the diagnosis and improved treatment of nervous system
XX diseases such as Alzheimer's disease, multiple sclerosis, schizophrenia,
XX epilepsy and cancers. The present sequence is a protein encoded by one of
XX the probes of the invention
XX Sequence 26 AA;
Query Match 34.0%; Score 33; DB 4; Length 26;

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Query Match

Best Local Similarity 43.8%; Pred. No. 2e+02; Mismatches 7; Conservative 2; Indels 7; Gaps 0;

Qy 3 NMLNSKIAFKIVSQEP 18  
 ||| :|||  
 Db 2 NTLEKTPQILGQEP 17

RESULT 9  
 ABG54477  
 ID ABG54477 standard; peptide; 26 AA.

XX AC ABG54477;  
 XX DT 25-FEB-2003 (first entry)  
 XX DE Human liver peptide, SEQ ID No 33125.  
 XX KW Human; liver; cirrhosis; hyperlipoproteinaemia; hyperlipidaemia;  
 KW hypercholesterolaemia; coronary heart disease.  
 XX OS Homo sapiens.

XX PN WO200157273-A2.  
 XX PD 09-AUG-2001.  
 XX PF 30-JAN-2001; 2001WO-US0000664.  
 XX PR 04-FEB-2000; 2000US-0180312P.  
 XX PR 26-MAY-2000; 2000US-0207456P.  
 XX PR 30-JUN-2000; 2000US-00608408.  
 XX PR 03-AUG-2000; 2000US-00632366.  
 XX PR 21-SEP-2000; 2000US-0234687P.  
 XX PR 27-SEP-2000; 2000US-0236359P.  
 XX PR 04-OCT-2000; 2000GB-00024263.  
 XX PA (MOLE-) MOLECULAR DYNAMICS INC.

XX PI Penn SG, Hanzel DK, Chen W, Rank DR;  
 XX WPI; 2001-488898/53.

XX PT Human genome-derived single exon nucleic acid probes useful for analyzing  
 PT gene expression in human adult liver.

XX PS Claim 27; SEQ ID NO 33125; 658pp; English.

XX CC The invention relates to a single exon nucleic acid probe (SENP) (I) for  
 CC measuring human gene expression in a sample derived from human adult  
 CC liver, comprising one of 13109 defined nucleotide sequences given in the  
 CC specification (or complements/ fragments). The probe hybridises at high  
 CC stringency to a nucleic acid molecule expressed in the human adult liver.  
 CC (I) may be used for predicting, measuring and displaying gene expression  
 CC in samples derived from human adult liver. The genes identified may be  
 CC involved in genetic liver diseases such as cirrhosis.  
 CC hyperlipoproteinaemia, hyperlipidaemia and hypercholesterolaemia which is  
 CC associated with coronary heart disease. ABG47348-ABG59930 represent human  
 CC liver single exon encoded peptides of the invention. Note: The sequence  
 CC information for this patent does not appear in the printed specification  
 CC but was obtained in electronic format directly from WIPO at  
 CC ftp.wipo.int/pub/published\_pct\_sequences

XX SQ Sequence 26 AA;

Query Match 34.0%; Score 33; DB 4; Length 26;  
 Best Local Similarity 43.8%; Pred. No. 2e+02;  
 Matches 7; Conservative 2; Mismatches 7; Indels 0; Gaps 0;

Qy 3 NMLNSKIAFKIVSQEP 18  
 ||| :|||  
 Db 2 NTLEKTPQILGQEP 17

RESULT 10  
 ABG42601

XX ID ABG42601 standard; peptide; 26 AA.

XX AC ABG42601;  
 XX DT 19-AUG-2002 (first entry)

XX DE Human peptide encoded by genome-derived single exon probe SEQ ID 32266.  
 XX KW Human; single exon probe; asthma; lung cancer; COPD; ILD;  
 KW chronic obstructive pulmonary disease; interstitial lung disease;  
 KW familial idiopathic pulmonary fibrosis; neurofibromatosis;  
 KW tuberous sclerosis; Gaucher's disease; Niemann-Pick disease;  
 KW Hermansky-Pudlak syndrome; sarcoidosis; pulmonary haemosiderosis;  
 KW pulmonary histiocytosis; lymphangiomyomatosis; Karagener syndrome;  
 KW pulmonary alveolar proteinosis; fibrocystic pulmonary dysplasia;  
 KW primary ciliary dyskinesia; pulmonary hypertension;  
 KW hyaline membrane disease.

XX OS Homo sapiens.  
 XX PN WO200186003-A2.  
 XX PD 15-NOV-2001.

XX PF 30-JAN-2001; 2001WO-US0000665.

XX PR 04-FEB-2000; 2000US-0180312P.  
 XX PR 26-MAY-2000; 2000US-0207456P.  
 XX PR 30-JUN-2000; 2000US-00608408.  
 XX PR 03-AUG-2000; 2000US-00632366.  
 XX PR 21-SEP-2000; 2000US-0234687P.  
 XX PR 27-SEP-2000; 2000US-0236359P.  
 XX PR 04-OCT-2000; 2000GB-00024263.  
 XX PA (MOLE-) MOLECULAR DYNAMICS INC.

XX PI Penn SG, Hanzel DK, Chen W, Rank DR;  
 XX WPI; 2002-114183/15.

XX PT Spatially-addressable set of single exon nucleic acid probes, used to  
 PT measure gene expression in human lung samples.

XX PS Claim 27; SEQ ID NO 32266; 634pp; English.

XX CC The invention relates to a spatially-addressable set of single exon  
 CC nucleic acid probes for measuring gene expression in a sample derived  
 CC from human lung comprising single exon nucleic acid probes having one of  
 CC 12614 nucleic acid sequences mentioned in the specification, or their  
 CC complements or the 12387 open reading frames derived from the 12614  
 CC probes. Also included are a microarray comprising the novel set of probes  
 CC ; the novel set of probes which hybridise at high stringency to a nucleic  
 CC acid expressed in the human lung; measuring gene expression in a sample  
 CC derived from human lung, comprising (a) contacting the array with a  
 CC collection of detectably labeled nucleic acids derived from human lung  
 CC mRNA, and (b) measuring the label detectably bound to each probe of the  
 CC array; identifying exons in a eukaryotic genome, comprising (a)  
 CC algorithmically predicting at least one exon from genomic sequences of  
 CC the eukaryote; and (b) detecting specific hybridisation of detectably  
 CC labeled nucleic acids from eukaryote lung mRNA, to a single exon probe,  
 CC having a fragment identical to the predicted exon, the probe is included  
 CC in the above mentioned microarray; assigning exons to a single gene,  
 CC comprising (a) identifying exons from genomic sequence by the method  
 CC above and (b) measuring the expression of each of the exons in several  
 CC tissues and/or cell types using hybridisation to a single exon  
 CC microarrays having a probe with the exon, where a common pattern of  
 CC expression of the exons in the tissues and/or cell types indicates that  
 CC the exons should be assigned to a single gene; a peptide comprising one  
 CC of 12011 sequences, mentioned in the specification, or encoded by the  
 CC probes/open reading frames (ORF). The probes are used for gene expression

CC analysis, and for identifying exons in a gene, particularly using human  
 CC lung derived mRNA and for the study of lung diseases such as asthma, lung  
 CC cancer, chronic obstructive pulmonary disease (COPD), interstitial lung  
 CC disease (ILD), familial idiopathic pulmonary fibrosis, neurofibromatosis,  
 CC tuberous sclerosis, Gaucher's disease, Niemann-Pick disease, Hermansky-  
 CC Pudlak syndrome, sarcoidosis, pulmonary haemosiderosis, pulmonary  
 CC histiocytosis, lymphangioleiomyomatosis, pulmonary alveolar proteinosis,  
 CC Karagenen syndrome, fibrocystic pulmonary dysplasia, primary ciliary  
 CC dyskinesia, pulmonary hypertension and hyaline membrane disease. The  
 CC present sequence is a peptide/protein encoded by a single exon probe of  
 CC the invention. Note: The sequence data for this patent did not form part  
 CC of the printed specification, but was obtained in electronic format  
 CC directly from WIFO at ftp.wipo.int/pub/published\_pct\_sequences  
 XX  
 SQ Sequence 26 AA;

Query Match 34.0%; Score 33; DB 5; Length 26;

Best Local Similarity 43.8%; Pred. No. 2e+02;

Matches 7; Conservative 2; Mismatches 7; Indels 0; Gaps 0;

QY 3 NHLNSKIAFKIVSQEP 18

Db 2 NTLERKTIQLGQEP 17

RESULT 11

AAG27280

ID AAG27280 standard; protein; 44 AA.

XX AAG27280;

XX 17-OCT-2000 (first entry)

XX Zea mays protein fragment SEQ ID NO: 32054.

XX Protein identification; signal transduction pathway; metabolic pathway;

KW hybridisation assay; genetic mapping; gene expression control; promoter;

KW termination sequence; corn.

XX Zea mays subsp. mays.

XX EP1033405-A2.

XX 06-SEP-2000.

XX 25-FEB-2000; 2000EP-00301439.

XX 25-FEB-1999; 99US-0121825P.

PR 05-MAR-1999; 99US-0123180P.

PR 09-MAR-1999; 99US-0123548P.

PR 23-MAR-1999; 99US-0125788P.

PR 25-MAR-1999; 99US-0126264P.

PR 29-MAR-1999; 99US-0126785P.

PR 01-APR-1999; 99US-0127462P.

PR 06-APR-1999; 99US-0128234P.

PR 08-APR-1999; 99US-0128714P.

PR 16-APR-1999; 99US-0129845P.

PR 19-APR-1999; 99US-0130077P.

PR 21-APR-1999; 99US-0130449P.

PR 23-APR-1999; 99US-0130510P.

PR 28-APR-1999; 99US-0130891P.

PR 30-APR-1999; 99US-0131449P.

PR 30-APR-1999; 99US-0132048P.

PR 04-MAY-1999; 99US-0132407P.

PR 05-MAY-1999; 99US-0132484P.

PR 06-MAY-1999; 99US-0132485P.

PR 06-MAY-1999; 99US-0132487P.

PR 07-MAY-1999; 99US-0132863P.

PR 11-MAY-1999; 99US-0134256P.

PR 14-MAY-1999; 99US-0134218P.

PR 14-MAY-1999; 99US-0134219P.

PR 14-MAY-1999; 99US-0134221P.

PR 14-MAY-1999; 99US-0134370P.

PR 18-MAY-1999; 99US-0134768P.

PR 19-MAY-1999; 99US-0134941P.

PR 20-MAY-1999; 99US-0135124P.

PR 21-MAY-1999; 99US-0135353P.

PR 24-MAY-1999; 99US-0135629P.

PR 25-MAY-1999; 99US-0136021P.

PR 27-MAY-1999; 99US-0136392P.

PR 28-MAY-1999; 99US-0136782P.

PR 01-JUN-1999; 99US-0137222P.

PR 03-JUN-1999; 99US-0137528P.

PR 04-JUN-1999; 99US-0137502P.

PR 07-JUN-1999; 99US-0137724P.

PR 08-JUN-1999; 99US-0138094P.

PR 10-JUN-1999; 99US-0138540P.

PR 10-JUN-1999; 99US-0138847P.

PR 14-JUN-1999; 99US-0139115P.

PR 16-JUN-1999; 99US-0139452P.

PR 16-JUN-1999; 99US-0139453P.

PR 17-JUN-1999; 99US-0139492P.

PR 18-JUN-1999; 99US-0139454P.

PR 18-JUN-1999; 99US-0139455P.

PR 18-JUN-1999; 99US-0139456P.

PR 18-JUN-1999; 99US-0139462P.

PR 18-JUN-1999; 99US-0139463P.

PR 18-JUN-1999; 99US-0139750P.

PR 18-JUN-1999; 99US-0139763P.

PR 21-JUN-1999; 99US-0139817P.

PR 22-JUN-1999; 99US-0139899P.

PR 23-JUN-1999; 99US-0140353P.

PR 24-JUN-1999; 99US-0140354P.

PR 24-JUN-1999; 99US-0140695P.

PR 28-JUN-1999; 99US-0140823P.

PR 29-JUN-1999; 99US-0140991P.

PR 30-JUN-1999; 99US-0141287P.

PR 01-JUL-1999; 99US-0141842P.

PR 01-JUL-1999; 99US-0142154P.

PR 02-JUL-1999; 99US-0142055P.

PR 06-JUL-1999; 99US-0142390P.

PR 08-JUL-1999; 99US-0142803P.

PR 09-JUL-1999; 99US-0142920P.

PR 12-JUL-1999; 99US-0142977P.

PR 13-JUL-1999; 99US-0143542P.

PR 14-JUL-1999; 99US-0143624P.

PR 15-JUL-1999; 99US-0144005P.

PR 16-JUL-1999; 99US-0144085P.

PR 16-JUL-1999; 99US-0144086P.

PR 19-JUL-1999; 99US-0144325P.

PR 19-JUL-1999; 99US-0144331P.

PR 19-JUL-1999; 99US-0144332P.

PR 19-JUL-1999; 99US-0144333P.

PR 19-JUL-1999; 99US-0144334P.

PR 19-JUL-1999; 99US-0144335P.

PR 20-JUL-1999; 99US-0144352P.

PR 20-JUL-1999; 99US-0144632P.

PR 20-JUL-1999; 99US-0144884P.

PR 21-JUL-1999; 99US-0144814P.

PR 21-JUL-1999; 99US-0145086P.

PR 21-JUL-1999; 99US-0145088P.

PR 22-JUL-1999; 99US-0145085P.

PR 22-JUL-1999; 99US-0145087P.

PR 22-JUL-1999; 99US-0145089P.

PR 22-JUL-1999; 99US-0145192P.

PR 23-JUL-1999; 99US-0145145P.

PR 23-JUL-1999; 99US-0145218P.

PR 23-JUL-1999; 99US-0145224P.

PR 26-JUL-1999; 99US-0145276P.

PR 27-JUL-1999; 99US-0145913P.

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PR 27-JUL-1999; 99US-0145918P.
PR 27-JUL-1999; 99US-0145919P.
PR 28-JUL-1999; 99US-0145951P.
PR 02-AUG-1999; 99US-0146386P.
PR 02-AUG-1999; 99US-0146388P.
PR 02-AUG-1999; 99US-0146389P.
PR 03-AUG-1999; 99US-0147038P.
PR 04-AUG-1999; 99US-0147204P.
PR 04-AUG-1999; 99US-0147302P.
PR 05-AUG-1999; 99US-0147192P.
PR 05-AUG-1999; 99US-0147260P.
PR 06-AUG-1999; 99US-0147303P.
PR 06-AUG-1999; 99US-0147416P.
PR 09-AUG-1999; 99US-0147493P.
PR 09-AUG-1999; 99US-0147935P.
PR 10-AUG-1999; 99US-0148171P.
PR 11-AUG-1999; 99US-0148319P.
PR 12-AUG-1999; 99US-0148341P.
PR 13-AUG-1999; 99US-0148565P.
PR 13-AUG-1999; 99US-0148684P.
PR 16-AUG-1999; 99US-0149368P.
PR 17-AUG-1999; 99US-0149175P.
PR 18-AUG-1999; 99US-0149426P.
PR 20-AUG-1999; 99US-0149722P.
PR 20-AUG-1999; 99US-0149723P.
PR 20-AUG-1999; 99US-0149929P.
PR 23-AUG-1999; 99US-0149930P.
PR 23-AUG-1999; 99US-0150566P.
PR 26-AUG-1999; 99US-0150884P.
PR 27-AUG-1999; 99US-0151065P.
PR 27-AUG-1999; 99US-0151066P.
PR 27-AUG-1999; 99US-0151080P.
PR 30-AUG-1999; 99US-0151303P.
PR 31-AUG-1999; 99US-0151438P.
PR 01-SEP-1999; 99US-0151930P.
PR 07-SEP-1999; 99US-0152363P.
PR 10-SEP-1999; 99US-0153070P.
PR 13-SEP-1999; 99US-0153758P.
PR 15-SEP-1999; 99US-0154018P.
PR 16-SEP-1999; 99US-0154039P.
PR 20-SEP-1999; 99US-0154779P.
PR 22-SEP-1999; 99US-0155139P.
PR 23-SEP-1999; 99US-0155486P.
PR 24-SEP-1999; 99US-0155659P.
PR 28-SEP-1999; 99US-0156458P.
PR 28-SEP-1999; 99US-0156596P.
PR 04-OCT-1999; 99US-0157117P.
PR 05-OCT-1999; 99US-0157753P.
PR 06-OCT-1999; 99US-0157865P.
PR 07-OCT-1999; 99US-0158029P.
PR 08-OCT-1999; 99US-0158232P.
PR 12-OCT-1999; 99US-0158369P.
PR 13-OCT-1999; 99US-0159293P.
PR 13-OCT-1999; 99US-0159294P.
PR 14-OCT-1999; 99US-0159329P.
PR 14-OCT-1999; 99US-0159330P.
PR 14-OCT-1999; 99US-0159331P.
PR 14-OCT-1999; 99US-0159637P.
PR 14-OCT-1999; 99US-0159638P.
PR 18-OCT-1999; 99US-0159584P.
PR 21-OCT-1999; 99US-0160741P.
PR 21-OCT-1999; 99US-0160767P.
PR 21-OCT-1999; 99US-0160768P.
PR 21-OCT-1999; 99US-0160770P.
PR 21-OCT-1999; 99US-0160814P.
PR 21-OCT-1999; 99US-0160815P.
PR 22-OCT-1999; 99US-0160980P.
PR 22-OCT-1999; 99US-0160981P.
PR 22-OCT-1999; 99US-0160989P.
PR 25-OCT-1999; 99US-0161404P.
PR 25-OCT-1999; 99US-0161405P.
PR 25-OCT-1999; 99US-0161406P.
PR 26-OCT-1999; 99US-0161359P.
PR 26-OCT-1999; 99US-0161360P.
PR 26-OCT-1999; 99US-0161361P.
PR 28-OCT-1999; 99US-0161920P.
PR 28-OCT-1999; 99US-0161922P.
PR 28-OCT-1999; 99US-0161993P.
PR 29-OCT-1999; 99US-0162142P.

Query Match 34.0%; Score 33; DB 3; Length 44;
Best Local Similarity 46.2%; Pred. No. 3.7e+02;
Matches 6; Conservative 4; Mismatches 3; Indels 0; Gaps 0;

Qy 1 EPNHLNLSKIAFKI 13
Db 10 QPNHLDSWGSFAL 22

:|||||:|:
:|||||:|:

RESULT 12
AAG59652
ID AAG59652 standard; protein; 36 AA.
XX AAG59652;
XX 18-OCT-2000 (first entry)
DE Arabidopsis thaliana protein fragment SEQ ID NO: 77180.
KW Protein identification; signal transduction pathway; metabolic pathway;
KW hybridisation assay; genetic mapping; gene expression control; promoter;
XX termination sequence.
OS Arabidopsis thaliana.
XX
PN EP1033405-A2.
XX
PD 06-SEP-2000.
XX
PF 25-FEB-2000; 2000EP-00301439.
XX
PR 25-FEB-1999; 99US-0121825P.
PR 05-MAR-1999; 99US-0123180P.
PR 09-MAR-1999; 99US-0123548P.
PR 23-MAR-1999; 99US-0125788P.
PR 25-MAR-1999; 99US-0126264P.
PR 23-MAR-1999; 99US-0126785P.
PR 01-APR-1999; 99US-0127462P.
PR 06-APR-1999; 99US-0128234P.
PR 08-APR-1999; 99US-0128714P.
PR 16-APR-1999; 99US-0129845P.
PR 19-APR-1999; 99US-0130077P.
PR 21-APR-1999; 99US-0130449P.
PR 23-APR-1999; 99US-0130510P.
PR 23-APR-1999; 99US-0130891P.
PR 28-APR-1999; 99US-0131449P.
PR 30-APR-1999; 99US-0132048P.
PR 30-APR-1999; 99US-0132407P.
PR 04-MAY-1999; 99US-0132484P.
PR 05-MAY-1999; 99US-0132485P.
PR 06-MAY-1999; 99US-0132486P.
PR 06-MAY-1999; 99US-0132487P.
PR 07-MAY-1999; 99US-0132863P.
PR 11-MAY-1999; 99US-0134256P.
PR 14-MAY-1999; 99US-0134218P.
PR 14-MAY-1999; 99US-0134219P.
PR 14-MAY-1999; 99US-0134221P.
PR 14-MAY-1999; 99US-0134370P.
PR 18-MAY-1999; 99US-0134768P.
PR 19-MAY-1999; 99US-0134941P.
PR 20-MAY-1999; 99US-0135124P.
PR 21-MAY-1999; 99US-0135353P.
PR 24-MAY-1999; 99US-0135629P.
PR 25-MAY-1999; 99US-0136021P.
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PR 28-MAY-1999; 99US-0136782P.  
PR 01-JUN-1999; 99US-0137222P.  
PR 03-JUN-1999; 99US-0137528P.  
PR 04-JUN-1999; 99US-0137502P.  
PR 07-JUN-1999; 99US-0137724P.  
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PR 10-JUN-1999; 99US-0138540P.  
PR 10-JUN-1999; 99US-0138847P.  
PR 14-JUN-1999; 99US-0139119P.  
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PR 17-JUN-1999; 99US-0139492P.  
PR 18-JUN-1999; 99US-0139454P.  
PR 18-JUN-1999; 99US-0139455P.  
PR 18-JUN-1999; 99US-0139458P.  
PR 18-JUN-1999; 99US-0139457P.  
PR 18-JUN-1999; 99US-0139458P.  
PR 18-JUN-1999; 99US-0139459P.  
PR 18-JUN-1999; 99US-0139459P.  
PR 18-JUN-1999; 99US-0139460P.  
PR 18-JUN-1999; 99US-0139461P.  
PR 18-JUN-1999; 99US-0139462P.  
PR 18-JUN-1999; 99US-0139463P.  
PR 18-JUN-1999; 99US-0139750P.  
PR 18-JUN-1999; 99US-0139763P.  
PR 21-JUN-1999; 99US-0139817P.  
PR 21-JUN-1999; 99US-0139899P.  
PR 23-JUN-1999; 99US-0140353P.  
PR 23-JUN-1999; 99US-0140354P.  
PR 24-JUN-1999; 99US-0140695P.  
PR 28-JUN-1999; 99US-0140823P.  
PR 29-JUN-1999; 99US-0140991P.  
PR 30-JUN-1999; 99US-0141287P.  
PR 01-JUL-1999; 99US-0141842P.  
PR 01-JUL-1999; 99US-0142154P.  
PR 02-JUL-1999; 99US-0142055P.  
PR 06-JUL-1999; 99US-0142390P.  
PR 08-JUL-1999; 99US-0142803P.  
PR 09-JUL-1999; 99US-0142920P.  
PR 12-JUL-1999; 99US-0142977P.  
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PR 14-JUL-1999; 99US-0143624P.  
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PR 16-JUL-1999; 99US-0144085P.  
PR 16-JUL-1999; 99US-0144086P.  
PR 19-JUL-1999; 99US-0144325P.  
PR 19-JUL-1999; 99US-0144331P.  
PR 19-JUL-1999; 99US-0144332P.  
PR 19-JUL-1999; 99US-0144333P.  
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PR 20-JUL-1999; 99US-0144352P.  
PR 20-JUL-1999; 99US-0144632P.  
PR 20-JUL-1999; 99US-0144684P.  
PR 21-JUL-1999; 99US-0144814P.  
PR 21-JUL-1999; 99US-0145086P.  
PR 21-JUL-1999; 99US-0145088P.  
PR 22-JUL-1999; 99US-0145089P.  
PR 22-JUL-1999; 99US-0145087P.  
PR 22-JUL-1999; 99US-0145089P.  
PR 22-JUL-1999; 99US-0145192P.  
PR 23-JUL-1999; 99US-0145214P.  
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PR 02-AUG-1999; 99US-0146389P.  
PR 03-AUG-1999; 99US-0147038P.

PR 04-AUG-1999; 99US-0147204P.  
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PR 05-AUG-1999; 99US-0147260P.  
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PR 06-AUG-1999; 99US-0147416P.  
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PR 12-AUG-1999; 99US-0148341P.  
PR 13-AUG-1999; 99US-0148565P.  
PR 13-AUG-1999; 99US-0148684P.  
PR 16-AUG-1999; 99US-0149368P.  
PR 17-AUG-1999; 99US-0149175P.  
PR 18-AUG-1999; 99US-0149426P.  
PR 20-AUG-1999; 99US-0149722P.  
PR 20-AUG-1999; 99US-0149723P.  
PR 20-AUG-1999; 99US-0149929P.  
PR 23-AUG-1999; 99US-0149902P.  
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PR 27-AUG-1999; 99US-0151066P.  
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PR 30-AUG-1999; 99US-0151303P.  
PR 31-AUG-1999; 99US-0151438P.  
PR 01-SEP-1999; 99US-0151930P.  
PR 07-SEP-1999; 99US-0152363P.  
PR 10-SEP-1999; 99US-0153070P.  
PR 13-SEP-1999; 99US-0153758P.  
PR 15-SEP-1999; 99US-0154018P.  
PR 16-SEP-1999; 99US-0154039P.  
PR 20-SEP-1999; 99US-0154779P.  
PR 22-SEP-1999; 99US-0155139P.  
PR 23-SEP-1999; 99US-0155486P.  
PR 24-SEP-1999; 99US-0155659P.  
PR 28-SEP-1999; 99US-0156458P.  
PR 29-SEP-1999; 99US-0156596P.  
PR 04-OCT-1999; 99US-0157117P.  
PR 05-OCT-1999; 99US-0157753P.  
PR 06-OCT-1999; 99US-0158029P.  
PR 07-OCT-1999; 99US-0158029P.  
PR 08-OCT-1999; 99US-0158232P.  
PR 12-OCT-1999; 99US-0158369P.  
PR 13-OCT-1999; 99US-0159293P.  
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PR 14-OCT-1999; 99US-0159330P.  
PR 14-OCT-1999; 99US-0159331P.  
PR 14-OCT-1999; 99US-0159637P.  
PR 14-OCT-1999; 99US-0159638P.  
PR 18-OCT-1999; 99US-0159584P.  
PR 21-OCT-1999; 99US-0160741P.  
PR 21-OCT-1999; 99US-0160767P.  
PR 21-OCT-1999; 99US-0160768P.  
PR 21-OCT-1999; 99US-0160770P.  
PR 21-OCT-1999; 99US-0160814P.  
PR 21-OCT-1999; 99US-0160815P.  
PR 22-OCT-1999; 99US-0160980P.  
PR 22-OCT-1999; 99US-0160981P.  
PR 22-OCT-1999; 99US-0160989P.  
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PR 25-OCT-1999; 99US-0161405P.  
PR 25-OCT-1999; 99US-0161406P.  
PR 26-OCT-1999; 99US-0161359P.  
PR 26-OCT-1999; 99US-0161360P.  
PR 26-OCT-1999; 99US-0161361P.  
PR 28-OCT-1999; 99US-0161920P.  
PR 28-OCT-1999; 99US-0161922P.  
PR 28-OCT-1999; 99US-0161993P.

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PR 29-OCT-1999; 99US-0162142P.
Query Match 33.5%; Score 32.5; DB 3; Length 36;
Best Local Similarity 57.1%; Pred. No. 3.6e+02;
Matches 8; Conservative 3; Mismatches 2; Indels 1; Gaps 1;

QY 3 NHLNSKIA-FKIVS 15
Db 14 HHLSLKITPFKVS 27

RESULT 13
AAM16072
ID AAM16072 standard; protein; 36 AA.
XX AC AAM16072;
XX DT 12-OCT-2001 (first entry)
XX DE Peptide #2506 encoded by probe for measuring cervical gene expression.
XX KW Probe; human; microarray; gene expression; cervical epithelial cell;
XX KW cervical cancer.
XX OS Homo sapiens.
XX PN WO200157278-A2.
XX PD 09-AUG-2001.
XX PF 30-JAN-2001; 2001WO-US000670.
XX PR 04-FEB-2000; 2000US-0180312P.
XX PR 26-MAY-2000; 2000US-0207456P.
XX PR 30-JUN-2000; 2000US-00608408.
XX PR 03-AUG-2000; 2000US-00632366.
XX PR 21-SEP-2000; 2000US-0234687P.
XX PR 27-SEP-2000; 2000US-0236359P.
XX PR 04-OCT-2000; 2000GB-00024263.
XX PA (MOLE-) MOLECULAR DYNAMICS INC.
XX PI Penn SG, Hanzel DK, Chen W, Rank DR;
XX WPI; 2001-488901/53.
XX PT Human genome-derived single exon nucleic acid probes useful for analyzing
XX gene expression in human cervical epithelial cells.
XX PS Claim 27; SEQ ID NO 20898; 487pp; English.
XX CC The present invention relates to human single exon nucleic acid probes
XX (SENP: see AAL10068-RA128459). The present sequence is a peptide encoded
XX by one such probe. The SENPs are derived from human Hela cells. The SENPs
XX can be used to produce a single exon microarray, which can be used for
XX measuring human gene expression in a sample derived from human cervical
XX epithelial cells. By measuring gene expression, the probes are therefore
XX useful in grading and/or staging of diseases of the cervix, notably
XX cervical cancer. Note: The sequence data for this patent did not form
XX part of the printed specification, but was obtained in electronic format
XX directly from WIPO at ftp.wipo.int/pub/published_pct_sequences
XX Sequence 36 AA;

Query Match 33.5%; Score 32.5; DB 4; Length 36;
Best Local Similarity 53.8%; Pred. No. 3.6e+02;
Matches 7; Conservative 2; Mismatches 3; Indels 1; Gaps 1;

QY 1 EPNHLSKIAFKI 13
Db 19 EPNH-NSLVVFL 30

RESULT 14
ABB35064
ID ABB35064 standard; peptide; 36 AA.
XX AC ABB35064;
XX DT 04-FEB-2002 (first entry)
XX DE Peptide #2570 encoded by human foetal liver single exon probe.
XX KW Human; foetal liver; gene expression; single exon nucleic acid probe.
XX OS Homo sapiens.
XX PN WO200157277-A2.
XX PD 09-AUG-2001.
XX PF 30-JAN-2001; 2001WO-US000669.
XX PR 04-FEB-2000; 2000US-0180312P.
XX PR 26-MAY-2000; 2000US-0207456P.
XX PR 30-JUN-2000; 2000US-00608408.
XX PR 03-AUG-2000; 2000US-00632366.
XX PR 21-SEP-2000; 2000US-0234687P.
XX PR 27-SEP-2000; 2000US-0236359P.
XX PR 04-OCT-2000; 2000GB-00024263.
XX PA (MOLE-) MOLECULAR DYNAMICS INC.
XX PI Penn SG, Hanzel DK, Chen W, Rank DR;
XX WPI; 2001-483447/52.
XX PT Human genome-derived single exon nucleic acid probes useful for analyzing
XX gene expression in human fetal liver.
XX PS Claim 27; SEQ ID NO 27699; 639pp + Sequence Listing; English.
XX CC The invention relates to a single exon nucleic acid probe for measuring
XX human gene expression in a sample derived from human foetal liver. The
XX single exon nucleic acid probes may be used for predicting, measuring and
XX displaying gene expression in samples derived from human fetal liver. The
XX present sequence is a peptide encoded by a single exon nucleic acid probe
XX of the invention. Note: The sequence data for this patent did not form
XX part of the printed specification, but was obtained in electronic format
XX directly from WIPO at ftp.wipo.int/pub/published_pct_sequences
XX Sequence 36 AA;

Query Match 33.5%; Score 32.5; DB 4; Length 36;
Best Local Similarity 53.8%; Pred. No. 3.6e+02;
Matches 7; Conservative 2; Mismatches 3; Indels 1; Gaps 1;

QY 1 EPNHLSKIAFKI 13
Db 19 EPNH-NSLVVFL 30

RESULT 15
AAM28566
ID AAM28566 standard; protein; 36 AA.
XX AC AAM28566;
XX DT 17-OCT-2001 (first entry)
XX DE Peptide #2603 encoded by probe for measuring placental gene expression.
XX KW Probe; microarray; human; placenta; antenatal diagnosis;
XX KW genetic disorder.
XX OS Homo sapiens.

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XX Example 4; SEQ ID NO 28554; 658pp + Sequence Listing; English.

PS The present invention provides a number of single exon nucleic acid

CC probes which are derived from genomic sequences expressed in the human

CC bone marrow. They can be used to measure gene expression in bone marrow

CC samples, which may enable the improved diagnosis and treatment of cancers

CC such as lymphoma, leukaemia and myeloma. The present sequence is a

CC protein encoded by one of the probes of the invention

XX

XX Sequence 36 AA;

XX

Query Match 33.5%; Score 32.5; DB 4; Length 36;

Best Local Similarity 53.8%; Pred. No. 3.6e+02;

Matches 7; Conservative 2; Mismatches 3; Indels 1; Gaps 1;

QY 1 EPNHLSKIAPKI 13

Db 19 EPNH-NSLLVFP 30

||||| : | :

RESULT 18

AM55878

ID AAM55878 standard; protein; 36 AA.

XX

XX AAM55878;

AC

XX

DT 05-NOV-2001 (first entry)

XX

DE Human brain expressed single exon probe encoded protein SEQ ID NO: 27983.

XX

XX Human; brain expressed exon; gene expression analysis; probe; microarray;

KW Alzheimer's disease; multiple sclerosis; schizophrenia; epilepsy; cancer.

XX

OS Homo sapiens.

XX

PN WO200157275-A2.

XX

PD 05-AUG-2001.

XX

PF 30-JAN-2001; 2001WO-US000667.

XX

PR 04-FEB-2000; 2000US-0180312P.

PR 26-MAY-2000; 2000US-0207456P.

PR 30-JUN-2000; 2000US-00608408.

PR 03-AUG-2000; 2000US-00632366.

PR 21-SEP-2000; 2000US-0234687P.

PR 27-SEP-2000; 2000US-0236359P.

PR 04-OCT-2000; 2000GB-00024263.

XX

PA (MOLE-) MOLECULAR DYNAMICS INC.

XX

PI Penn SG, Hanzel DK, Chen W, Rank DR;

XX

XX WPI; 2001-483446/52.

XX

DR Single exon nucleic acid probes for analyzing gene expression in human

XX brains.

XX

XX Example 4; SEQ ID NO 27983; 650pp + Sequence Listing; English.

XX

XX The present invention provides a number of single exon nucleic acid

CC probes which are derived from genomic sequences expressed in the human

CC brain. They can be used to measure gene expression in brain cell samples,

CC which may enable the diagnosis and improved treatment of nervous system

CC diseases such as Alzheimer's disease, multiple sclerosis, schizophrenia,

CC epilepsy and cancers. The present sequence is a protein encoded by one of

CC the probes of the invention

XX

XX Sequence 36 AA;

XX

Query Match 33.5%; Score 32.5; DB 4; Length 36;

Best Local Similarity 53.8%; Pred. No. 3.6e+02;

Matches 7; Conservative 2; Mismatches 3; Indels 1; Gaps 1;

QY 1 EPNHLSKIAPKI 13

Db 19 EPNH-NSLLVFP 30

||||| : | :

Matches 7; Conservative 2; Mismatches 3; Indels 1; Gaps 1;

QY 1 EPNHLSKIAPKI 13

Db 19 EPNH-NSLLVFP 30

||||| : | :

RESULT 19

ABG49902

ID ABG49902 standard; peptide; 36 AA.

XX

XX ABG49902;

AC

XX

DT 25-FEB-2003 (first entry)

XX

DE Human liver peptide, SEQ ID No 28550.

XX

XX Human; liver; cirrhosis; hyperlipoproteinaemia; hyperlipidaemia;

KW hypercholesterolaemia; coronary heart disease.

XX

OS Homo sapiens.

XX

PN WO200157273-A2.

XX

PD 09-AUG-2001.

XX

PF 30-JAN-2001; 2001WO-US000664.

XX

PR 04-FEB-2000; 2000US-0180312P.

PR 26-MAY-2000; 2000US-0207456P.

PR 30-JUN-2000; 2000US-00608408.

PR 03-AUG-2000; 2000US-00632366.

PR 21-SEP-2000; 2000US-0234687P.

PR 27-SEP-2000; 2000US-0236359P.

PR 04-OCT-2000; 2000GB-00024263.

XX

PA (MOLE-) MOLECULAR DYNAMICS INC.

XX

PI Penn SG, Hanzel DK, Chen W, Rank DR;

XX

XX WPI; 2001-488898/53.

XX

DR Human genome-derived single exon nucleic acid probes useful for analyzing

XX gene expression in human adult liver.

XX

PS Claim 27; SEQ ID NO 28550; 658pp; English.

XX

CC The invention relates to a single exon nucleic acid probe (SENP) (I) for

CC measuring human gene expression in a sample derived from human adult

CC liver, comprising one of 13109 defined nucleotide sequences given in the

CC specification (or complements/fragments). The probe hybridises at high

CC stringency to a nucleic acid molecule expressed in the human adult liver.

CC (I) may be used for predicting, measuring and displaying gene expression

CC in samples derived from human adult liver. The genes identified may be

CC involved in genetic liver diseases such as cirrhosis,

CC hyperlipoproteinaemia, hyperlipidaemia and hypercholesterolaemia which is

CC associated with coronary heart disease. ABG47348-ABG5930 represent human

CC liver single exon encoded peptides of the invention. Note: The sequence

CC information for this patent does not appear in the printed specification

CC but was obtained in electronic format directly from WIPO at

CC ftp.wipo.int/pub/published\_pct\_sequences

XX

XX Sequence 36 AA;

XX

Query Match 33.5%; Score 32.5; DB 4; Length 36;

Best Local Similarity 53.8%; Pred. No. 3.6e+02;

Matches 7; Conservative 2; Mismatches 3; Indels 1; Gaps 1;

QY 1 EPNHLSKIAPKI 13

Db 19 EPNH-NSLLVFP 30

||||| : | :



CC present sequence is a peptide/protein encoded by a single exon probe of  
 CC the invention. Note: The sequence data for this patent did not form part  
 CC of the printed specification, but was obtained in electronic format  
 CC directly from WIPO at ftp.wipo.int/pub/published\_pct\_sequences

SQ Sequence 36 AA;

Query Match 33.5%; Score 32.5; DB 5; Length 36;  
 Best Local Similarity 53.8%; Pred. No. 3.6e+02;  
 Matches 7; Conservative 2; Mismatches 3; Indels 1; Gaps 1;

Qy 1 EPNHLSKIAFKI 13  
 ||||| : : :  
 Db 19 EPNH-NSLLVPL 30

RESULT 22  
 ABO12904  
 ID ABO12904 standard; peptide; 24 AA.  
 XX AC ABO12904;

DT 25-AUG-2003 (first entry)

XX Mouse zinc finger DNA binding domain #10.

DE Composite binding polypeptide; zinc finger nucleic acid binding domain;  
 XX autoimmune disorder; immunosuppressive; zinc finger DNA binding domain;  
 KW mouse.

XX Mus sp.

XX WO200299084-A2.

XX 12-DEC-2002.

XX 04-APR-2002; 2002WO-US022272.

XX 04-APR-2001; 2001GB-00008491.

XX (SANG-) SANGAMO BIOSCIENCES INC.

PI Moore M, Sepp A, Isalan M, Choo Y;

XX WPI; 2003-278214/27.

XX New composite binding zinc finger polypeptide, useful for designing  
 PT sequence-specific binding proteins regulating gene expression in the  
 PT fields of molecular biology, and for the diagnosis and treatment of  
 PT autoimmune disorders.

XX Example 3; Page 97; 157pp; English.

CC The invention relates to a composite binding polypeptide comprising a  
 CC first natural binding domain derived from a first natural binding  
 CC polypeptide and a second natural binding domain derived from a second  
 CC natural binding polypeptide, where the first and second natural binding  
 CC polypeptides may be the same or different and where the polypeptide binds  
 CC to a target differing from the natural target of both the first and  
 CC second binding polypeptides. The invention also relates to a chimeric  
 CC polypeptide comprising a binding polypeptide cited above and a biological  
 CC effector domain, a library of natural binding domains, a library of  
 CC natural zinc finger nucleic acid binding domains comprising a linker  
 CC attached to it, a method for selecting a binding polypeptide capable of  
 CC binding to a target site and a method for designing a composite binding  
 CC polypeptide. The methods and compositions of the present invention are  
 CC useful for designing sequence-specific binding proteins for regulation of  
 CC gene expression in the fields of molecular biology. They can also be used  
 CC for the diagnosis and treatment of autoimmune disorders, and as research  
 CC tools and in transgenic animals. This sequence represents a mouse zinc  
 CC finger DNA binding domain used in the scope of the invention

XX Sequence 24 AA;

Query Match 33.0%; Score 32; DB 6; Length 24;  
 Best Local Similarity 62.5%; Pred. No. 2.7e+02;  
 Matches 5; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

Qy 2 PNHLNSKI 9  
 | : ||| :  
 Db 13 PDHLNSHV 20

RESULT 23  
 ABO11938  
 ID ABO11938 standard; peptide; 24 AA.  
 XX AC ABO11938;

DT 25-AUG-2003 (first entry)

XX Human zinc finger DNA binding domain #238.

DE Composite binding polypeptide; zinc finger nucleic acid binding domain;  
 XX autoimmune disorder; immunosuppressive; zinc finger DNA binding domain;  
 KW human.

XX Homo sapiens.

XX WO200299084-A2.

XX 12-DEC-2002.

XX 04-APR-2002; 2002WO-US022272.

XX 04-APR-2001; 2001GB-00008491.

XX (SANG-) SANGAMO BIOSCIENCES INC.

PI Moore M, Sepp A, Isalan M, Choo Y;

XX WPI; 2003-278214/27.

XX New composite binding zinc finger polypeptide, useful for designing  
 PT sequence-specific binding proteins regulating gene expression in the  
 PT fields of molecular biology, and for the diagnosis and treatment of  
 PT autoimmune disorders.

XX Example 2; Page 76; 157pp; English.

CC The invention relates to a composite binding polypeptide comprising a  
 CC first natural binding domain derived from a first natural binding  
 CC polypeptide and a second natural binding domain derived from a second  
 CC natural binding polypeptide, where the first and second natural binding  
 CC polypeptides may be the same or different and where the polypeptide binds  
 CC to a target differing from the natural target of both the first and  
 CC second binding polypeptides. The invention also relates to a chimeric  
 CC polypeptide comprising a binding polypeptide cited above and a biological  
 CC effector domain, a library of natural binding domains, a library of  
 CC natural zinc finger nucleic acid binding domains comprising a linker  
 CC attached to it, a method for selecting a binding polypeptide capable of  
 CC binding to a target site and a method for designing a composite binding  
 CC polypeptide. The methods and compositions of the present invention are  
 CC useful for designing sequence-specific binding proteins for regulation of  
 CC gene expression in the fields of molecular biology. They can also be used  
 CC for the diagnosis and treatment of autoimmune disorders, and as research  
 CC tools and in transgenic animals. This sequence represents a human zinc  
 CC finger DNA binding domain used in the scope of the invention

XX Sequence 24 AA;

Query Match 33.0%; Score 32; DB 6; Length 24;  
 Best Local Similarity 62.5%; Pred. No. 2.7e+02;  
 Matches 5; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

Qy 2 PNHLNSKI 9

Db 13 PDHLNSHV 20

RESULT 24  
ABU62168

ID ABU62168 standard; peptide; 24 AA.

AC ABU62168;

XX 25-AUG-2003 (first entry)

DT Human zinc finger DNA binding domain #6.

DE Composite binding polypeptide; zinc finger nucleic acid binding domain;

XX autoimmune disorder; immunosuppressive; zinc finger DNA binding domain;

KW human.

XX Homo sapiens.

OS WO200299084-A2.

EN 12-DEC-2002.

XX 04-APR-2002; 2002WO-US022272.

PF 04-APR-2001; 2001GB-00008491.

XX (SANG-) SANGAMO BIOSCIENCES INC.

FA Moore M, Sepp A, Isalan M, Choo Y;

PI WPI; 2003-278214/27.

DR New composite binding zinc finger polypeptide, useful for designing

XX sequence-specific binding proteins regulating gene expression in the

PT fields of molecular biology, and for the diagnosis and treatment of

PT autoimmune disorders.

XX Example 1; Page 70; 157pp; English.

PS The invention relates to a composite binding polypeptide comprising a

CC first natural binding domain derived from a first natural binding

CC polypeptide and a second natural binding domain derived from a second

CC natural binding polypeptide, where the first and second natural binding

CC polypeptides may be the same or different and where the polypeptide binds

CC to a target differing from the natural target of both the first and

CC second binding polypeptides. The invention also relates to a chimeric

CC polypeptide comprising a binding polypeptide cited above and a biological

CC effector domain, a library of natural binding domains, a library of

CC natural zinc finger nucleic acid binding domains comprising a linker

CC attached to it, a method for selecting a binding polypeptide capable of

CC binding to a target site and a method for designing a composite binding

CC polypeptide. The methods and compositions of the present invention are

CC useful for designing sequence-specific binding proteins for regulation of

CC gene expression in the fields of molecular biology. They can also be used

CC for the diagnosis and treatment of autoimmune disorders, and as research

CC tools and in transgenic animals. This sequence represents a human zinc

CC finger DNA binding domain used in the scope of the invention

XX Sequence 24 AA;

QY Query Match 33.0%; Score 32; DB 6; Length 24;  
Best Local Similarity 62.5%; Pred. No. 2.7e+02;  
Matches 5; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

Db 2 PNHLNSKI 9  
13 PDHLNSHV 20

RESULT 25  
AAW60192

ID AAW60192 standard; protein; 26 AA.

AC AAW60192;

XX 27-AUG-2003 (revised)

DT 03-SEP-1998 (first entry)

XX Bacteriophage spo1 Pol I-type DNA polymerase helix O region sequence.

DE Pol I-type; non-discriminating DNA polymerase; leprosy; tuberculosis;

KW mycobacteria; Bacteriophage spo1.

XX Bacteriophage SPO1.

OS US5776673-A.

PN 07-JUL-1998.

XX 21-APR-1995; 95US-00427072.

PF 21-APR-1995; 95US-00427072.

PR (HARD ) HARVARD COLLEGE.

FA Tabor S, Richardson CC;

PI WPI; 1998-398014/34.

DR Detecting mycobacteria for diagnosing tuberculosis and/or leprosy -

XX comprises detecting the presence of a DNA polymerase that does not

PT discriminate between deoxy- and di:deoxy-nucleoside triphosphate(s).

PT Disclosure; Col 5; 21pp; English.

PS This sequence represents the helix O region of Bacteriophage spo1 Pol I-

CC type DNA polymerase. This can be used in the method of invention of

CC diagnosing the presence of a mycobacterial organism. The method comprises

CC providing a sample from a patient, detecting the presence of a non-

CC discriminating DNA polymerase, not normally present by determining the

CC ability of the non-discriminating DNA polymerase to incorporate

CC dideoxynucleotides (ddNTP) relative to deoxynucleotides (dNTP). The

CC presence of non-discriminating DNA polymerase is indicative of presence

CC of a mycobacterial organism. The method is used to diagnose tuberculosis

CC and leprosy. (Updated on 27-AUG-2003 to correct OS field.)

XX Sequence 26 AA;

QY Query Match 33.0%; Score 32; DB 2; Length 26;  
Best Local Similarity 57.1%; Pred. No. 3e+02;  
Matches 8; Conservative 1; Mismatches 5; Indels 0; Gaps 0;

Db 6 NSKIAPKIVSORPA 19  
6 SKKIQFGIVYORSA 19

RESULT 26  
ABB17107

ID ABB17107 standard; protein; 33 AA.

XX ABB17107;

XX 23-JAN-2002 (first entry)

DT Human nervous system related polypeptide SEQ ID NO 5764.

XX Human; nontropic; neuroprotective; cytostatic; dermatological; virucide;

KW immunosuppressive; antiinflammatory; anti-HIV; antibacterial; vulnery;

KW antiparkinsonian; antiskilling; antianaemic; antiarthritis; cancer;

KW antirheumatic; hepatotropic; cerebroprotective; antiinflammatory;

KW antiallergic; antidiabetic; antitumor; anticonvulsant; antifungal;

KW antiparasitic; cardiac; immune disorder; cardiovascular disorder;

KW neurological disease; infection; nephrotropic; gene therapy; vaccine.



XX OS Homo sapiens.  
XX PN WO200159063-A2.  
XX XX  
XX PD 16-AUG-2001.  
XX PF  
XX PF  
XX 17-JAN-2001; 2001WO-US001334.  
XX  
XX 31-JAN-2000; 2000US-0179065P.  
XX PR 04-FEB-2000; 2000US-0180628P.  
XX PR 24-FEB-2000; 2000US-0184664P.  
XX PR 02-MAR-2000; 2000US-0186350P.  
XX PR 16-MAR-2000; 2000US-0189874P.  
XX PR 17-MAR-2000; 2000US-0190076P.  
XX PR 18-APR-2000; 2000US-0198123P.  
XX PR 19-MAY-2000; 2000US-0205515P.  
XX PR 07-JUN-2000; 2000US-0209467P.  
XX PR 28-JUN-2000; 2000US-0214886P.  
XX PR 30-JUN-2000; 2000US-0215135P.  
XX PR 07-JUL-2000; 2000US-0216647P.  
XX PR 07-JUL-2000; 2000US-0216880P.  
XX PR 11-JUL-2000; 2000US-0217487P.  
XX PR 11-JUL-2000; 2000US-0217496P.  
XX PR 14-JUL-2000; 2000US-0218290P.  
XX PR 26-JUL-2000; 2000US-0220963P.  
XX PR 26-JUL-2000; 2000US-0220964P.  
XX PR 14-AUG-2000; 2000US-0224518P.  
XX PR 14-AUG-2000; 2000US-0224519P.  
XX PR 14-AUG-2000; 2000US-0225213P.  
XX PR 14-AUG-2000; 2000US-0225214P.  
XX PR 14-AUG-2000; 2000US-0225266P.  
XX PR 14-AUG-2000; 2000US-0225267P.  
XX PR 14-AUG-2000; 2000US-0225268P.  
XX PR 14-AUG-2000; 2000US-0225270P.  
XX PR 14-AUG-2000; 2000US-0225447P.  
XX PR 14-AUG-2000; 2000US-0225757P.  
XX PR 14-AUG-2000; 2000US-0225758P.  
XX PR 14-AUG-2000; 2000US-0225759P.  
XX PR 18-AUG-2000; 2000US-0226279P.  
XX PR 22-AUG-2000; 2000US-0226681P.  
XX PR 22-AUG-2000; 2000US-0226868P.  
XX PR 22-AUG-2000; 2000US-0227182P.  
XX PR 23-AUG-2000; 2000US-0227009P.  
XX PR 30-AUG-2000; 2000US-0228924P.  
XX PR 01-SEP-2000; 2000US-0229287P.  
XX PR 01-SEP-2000; 2000US-0229343P.  
XX PR 01-SEP-2000; 2000US-0229344P.  
XX PR 01-SEP-2000; 2000US-0229345P.  
XX PR 05-SEP-2000; 2000US-0229509P.  
XX PR 06-SEP-2000; 2000US-0229513P.  
XX PR 06-SEP-2000; 2000US-0230437P.  
XX PR 08-SEP-2000; 2000US-0230438P.  
XX PR 08-SEP-2000; 2000US-0231242P.  
XX PR 08-SEP-2000; 2000US-0231243P.  
XX PR 08-SEP-2000; 2000US-0231244P.  
XX PR 08-SEP-2000; 2000US-0231413P.  
XX PR 08-SEP-2000; 2000US-0231414P.  
XX PR 08-SEP-2000; 2000US-0232080P.  
XX PR 08-SEP-2000; 2000US-0232081P.  
XX PR 12-SEP-2000; 2000US-0231968P.  
XX PR 14-SEP-2000; 2000US-0232398P.  
XX PR 14-SEP-2000; 2000US-0232399P.  
XX PR 14-SEP-2000; 2000US-0232400P.  
XX PR 14-SEP-2000; 2000US-0232401P.  
XX PR 14-SEP-2000; 2000US-0233063P.  
XX PR 14-SEP-2000; 2000US-0233064P.  
XX PR 21-SEP-2000; 2000US-0233065P.  
XX PR 21-SEP-2000; 2000US-0234223P.  
XX PR 25-SEP-2000; 2000US-0234274P.  
XX PR 25-SEP-2000; 2000US-0234997P.  
XX PR 25-SEP-2000; 2000US-0234998P.  
XX  
PR 26-SEP-2000; 2000US-0235484P.  
PR 27-SEP-2000; 2000US-0235834P.  
PR 27-SEP-2000; 2000US-0235836P.  
PR 29-SEP-2000; 2000US-0236327P.  
PR 29-SEP-2000; 2000US-0236367P.  
PR 29-SEP-2000; 2000US-0236368P.  
PR 29-SEP-2000; 2000US-0236369P.  
PR 29-SEP-2000; 2000US-0236370P.  
PR 02-OCT-2000; 2000US-0236802P.  
PR 02-OCT-2000; 2000US-0237037P.  
PR 02-OCT-2000; 2000US-0237038P.  
PR 02-OCT-2000; 2000US-0237039P.  
PR 02-OCT-2000; 2000US-0237040P.  
PR 13-OCT-2000; 2000US-0239335P.  
PR 13-OCT-2000; 2000US-0239337P.  
PR 20-OCT-2000; 2000US-0240960P.  
PR 20-OCT-2000; 2000US-0241785P.  
PR 20-OCT-2000; 2000US-0241786P.  
PR 20-OCT-2000; 2000US-0241787P.  
PR 20-OCT-2000; 2000US-0241808P.  
PR 20-OCT-2000; 2000US-0241809P.  
PR 20-OCT-2000; 2000US-0241826P.  
PR 01-NOV-2000; 2000US-0242221P.  
PR 08-NOV-2000; 2000US-0244617P.  
PR 08-NOV-2000; 2000US-0244647P.  
PR 08-NOV-2000; 2000US-0246475P.  
PR 08-NOV-2000; 2000US-0246476P.  
PR 08-NOV-2000; 2000US-0246477P.  
PR 08-NOV-2000; 2000US-0246478P.  
PR 08-NOV-2000; 2000US-0246523P.  
PR 08-NOV-2000; 2000US-0246524P.  
PR 08-NOV-2000; 2000US-0246525P.  
PR 08-NOV-2000; 2000US-0246526P.  
PR 08-NOV-2000; 2000US-0246527P.  
PR 08-NOV-2000; 2000US-0246528P.  
PR 08-NOV-2000; 2000US-0246532P.  
PR 08-NOV-2000; 2000US-0246609P.  
PR 08-NOV-2000; 2000US-0246610P.  
PR 08-NOV-2000; 2000US-0246611P.  
PR 17-NOV-2000; 2000US-0249207P.  
PR 17-NOV-2000; 2000US-0249208P.  
PR 17-NOV-2000; 2000US-0249209P.  
PR 17-NOV-2000; 2000US-0249210P.  
PR 17-NOV-2000; 2000US-0249211P.  
PR 17-NOV-2000; 2000US-0249212P.  
PR 17-NOV-2000; 2000US-0249213P.  
PR 17-NOV-2000; 2000US-0249214P.  
PR 17-NOV-2000; 2000US-0249215P.  
PR 17-NOV-2000; 2000US-0249216P.  
PR 17-NOV-2000; 2000US-0249217P.  
PR 17-NOV-2000; 2000US-0249218P.  
PR 17-NOV-2000; 2000US-0249244P.  
PR 17-NOV-2000; 2000US-0249245P.  
PR 17-NOV-2000; 2000US-0249264P.  
PR 17-NOV-2000; 2000US-0249265P.  
PR 17-NOV-2000; 2000US-0249297P.  
PR 17-NOV-2000; 2000US-0249299P.  
PR 17-NOV-2000; 2000US-0249300P.  
PR 01-DEC-2000; 2000US-0250391P.  
PR 05-DEC-2000; 2000US-0251160P.  
PR 05-DEC-2000; 2000US-0251030P.  
PR 05-DEC-2000; 2000US-0251988P.  
PR 06-DEC-2000; 2000US-0256719P.  
PR 08-DEC-2000; 2000US-0251479P.  
PR 08-DEC-2000; 2000US-0251856P.  
PR 08-DEC-2000; 2000US-0251868P.  
PR 08-DEC-2000; 2000US-0251869P.  
PR 08-DEC-2000; 2000US-0251990P.  
PR 11-DEC-2000; 2000US-0254097P.  
PR 05-JAN-2001; 2001US-0259678P.  
XX XX

PA (HUMA-) HUMAN GENOME SCI INC.  
XX Rosen CA, Barash SC, Ruben SM;  
PI WPI; 2001-541565/60.  
XX N-PSDB; ABA13433.  
DR Nucleic acids encoding 3224 human nervous system antigen polypeptides,  
PT useful for preventing, diagnosing and/or treating nervous system cancers  
PT and metastases.  
XX Claim 11; SEQ ID NO 5764; 1701pp + Sequence Listing; English.  
XX  
XX The invention relates to novel genes (ABA11004-ABA21534) and proteins  
CC (ABA14678-ABA18001) useful for preventing, treating or ameliorating  
CC medical conditions e.g. by protein or gene therapy. The genes are  
CC isolated from a range of human tissues disclosed in the specification.  
CC The nucleic acids, proteins, antibodies and (ant)agonists are useful in  
CC the diagnosis, treatment and prevention of: (a) cancer, e.g. breast and  
CC ovarian cancer and other cancers of the adrenal gland, bone, bone marrow,  
CC breast, gastrointestinal tract, liver, lung, or urogenital; (b) immune  
CC disorders e.g. Addison's disease, allergies, autoimmune haemolytic  
CC anaemia, autoimmune thyroiditis, diabetes mellitus, Crohn's disease,  
CC multiple sclerosis, rheumatoid arthritis and ulcerative colitis; (c)  
CC cardiovascular disorders such as myocardial ischaemias; (d) wound healing  
CC ; (e) neurological diseases e.g. cerebral anoxia and epilepsy; and (f)  
CC infectious diseases such as viral, bacterial, fungal and parasitic  
CC infections. Note: The sequence data for this patent did not form part of  
CC the printed specification, but was obtained in electronic format directly  
CC from WIPO at ftp.wipo.int/pub/published\_pct\_sequences  
XX  
XX Sequence 33 AA;  
Query Match 33.0%; Score 32; DB 4; Length 33;  
Best Local Similarity 66.7%; Pred. No. 4e+02;  
Matches 6; Conservative 0; Mismatches 3; Indels 0; Gaps 0;  
QY 3 NHLNSKIAF 11  
DB 22 NHLNVSITF 30  
RESULT 27  
ABBL6071  
ID ABBL6071 standard; protein; 38 AA.  
XX  
XX ABBL6071;  
XX  
XX 23-JAN-2002 (first entry)  
XX  
XX Human nervous system related polypeptide SEQ ID NO 4728.  
XX  
XX Human; nootropic; neuroprotective; cytostatic; dermatological; virucide;  
KW immunosuppressive; antiinflammatory; anti-HIV; antibacterial; vulnary;  
KW antiparkinsonian; antiskilling; antianaemic; antiarthritic; cancer;  
KW antirheumatic; hepatotropic; cerebroprotective; antiinflammatory;  
KW antiallergic; antidiabetic; antiulcer; anticonvulsant; antifungal;  
KW antiparasitic; cardiac; immune disorder; cardiovascular disorder;  
KW neurological disease; infection; nephrotropic; gene therapy; vaccine.  
XX  
XX Homo sapiens.  
XX  
XX WO200159063-A2.  
XX  
XX 16-AUG-2001.  
XX  
XX 17-JAN-2001; 2001WO-US001334.  
XX  
XX 31-JAN-2000; 2000US-0179065P.  
PR 04-FEB-2000; 2000US-0180628P.  
PR 24-FEB-2000; 2000US-0184664P.  
PR 02-MAR-2000; 2000US-0186350P.  
PR 16-MAR-2000; 2000US-0189874P.  
PR  
PR 17-MAR-2000; 2000US-0190076P.  
PR 18-APR-2000; 2000US-0198123P.  
PR 19-MAY-2000; 2000US-0205515P.  
PR 07-JUN-2000; 2000US-0209467P.  
PR 28-JUN-2000; 2000US-0214886P.  
PR 30-JUN-2000; 2000US-0215135P.  
PR 07-JUL-2000; 2000US-0216647P.  
PR 07-JUL-2000; 2000US-0216880P.  
PR 11-JUL-2000; 2000US-0217487P.  
PR 11-JUL-2000; 2000US-0217496P.  
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PR 26-JUL-2000; 2000US-0220963P.  
PR 26-JUL-2000; 2000US-0220964P.  
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PR 23-AUG-2000; 2000US-0227009P.  
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PR 01-SEP-2000; 2000US-0229287P.  
PR 01-SEP-2000; 2000US-0229343P.  
PR 01-SEP-2000; 2000US-0229344P.  
PR 01-SEP-2000; 2000US-0229345P.  
PR 05-SEP-2000; 2000US-0229509P.  
PR 05-SEP-2000; 2000US-0229513P.  
PR 06-SEP-2000; 2000US-0230437P.  
PR 06-SEP-2000; 2000US-0230438P.  
PR 08-SEP-2000; 2000US-0231242P.  
PR 08-SEP-2000; 2000US-0231243P.  
PR 08-SEP-2000; 2000US-0231244P.  
PR 08-SEP-2000; 2000US-0231413P.  
PR 08-SEP-2000; 2000US-0231414P.  
PR 08-SEP-2000; 2000US-0232080P.  
PR 08-SEP-2000; 2000US-0232081P.  
PR 12-SEP-2000; 2000US-0231968P.  
PR 14-SEP-2000; 2000US-0232397P.  
PR 14-SEP-2000; 2000US-0232398P.  
PR 14-SEP-2000; 2000US-0232399P.  
PR 14-SEP-2000; 2000US-0232400P.  
PR 14-SEP-2000; 2000US-0232401P.  
PR 14-SEP-2000; 2000US-0233063P.  
PR 14-SEP-2000; 2000US-0233084P.  
PR 21-SEP-2000; 2000US-0233085P.  
PR 21-SEP-2000; 2000US-0234223P.  
PR 21-SEP-2000; 2000US-0234274P.  
PR 25-SEP-2000; 2000US-0234997P.  
PR 25-SEP-2000; 2000US-0234998P.  
PR 26-SEP-2000; 2000US-0235484P.  
PR 27-SEP-2000; 2000US-0235834P.  
PR 27-SEP-2000; 2000US-0235836P.  
PR 29-SEP-2000; 2000US-0236327P.  
PR 29-SEP-2000; 2000US-0236367P.  
PR 29-SEP-2000; 2000US-0236368P.  
PR 29-SEP-2000; 2000US-0236369P.  
PR 29-SEP-2000; 2000US-0236370P.  
PR 02-OCT-2000; 2000US-0237037P.  
PR 02-OCT-2000; 2000US-0237038P.  
PR 02-OCT-2000; 2000US-0237039P.  
PR 02-OCT-2000; 2000US-0237040P.  
PR 13-OCT-2000; 2000US-0239935P.  
PR



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Query Match          33.0%; Score 32; DB 3; Length 46;
Best Local Similarity 35.7%; Pred. No. 6e+02;
Matches 5; Conservative 6; Mismatches 3; Indels 0; Gaps 0;

QY 5 LNSKIAFKIVSQEP 18
   | : : : : :
Db 1 LDRGVLYRMNQEP 14

RESULT 29
AAM13846
ID AAM13846 standard; protein; 48 AA.
XX
AC AAM13846;
XX
XX 12-OCT-2001 (first entry)
XX
DE Peptide #280 encoded by probe for measuring cervical gene expression.
XX
KW Probe; human; microarray; gene expression; cervical epithelial cell;
KW cervical cancer.
XX
OS Homo sapiens.
XX
PN WO200157278-A2.
XX
PD 09-AUG-2001.
XX
PF 30-JAN-2001; 2001WO-US000670.
XX
PR 04-FEB-2000; 2000US-0180312P.
PR 26-MAY-2000; 2000US-0207456P.
PR 30-JUN-2000; 2000US-00608408.
PR 03-AUG-2000; 2000US-00632366.
PR 21-SEP-2000; 2000US-0234687P.
PR 27-SEP-2000; 2000US-0236359P.
PR 04-OCT-2000; 2000GB-00024263.
XX
PA (MOLE-) MOLECULAR DYNAMICS INC.
XX
PI Penn SG, Hanzel DK, Chen W, Rank DR;
XX
XX WPI; 2001-488901/53.
XX
XX Human genome-derived single exon nucleic acid probes useful for analyzing
XX gene expression in human cervical epithelial cells.
XX
PS Claim 27; SEQ ID NO 18672; 487pp; English.
XX
CC The present invention relates to human single exon nucleic acid probes
CC (SENP: see AAI10068-AA128459). The present sequence is a peptide encoded
CC by one such probe. The SENPs are derived from human HeLa cells. The SENPs
CC can be used to produce a single exon microarray, which can be used for
CC measuring human gene expression in a sample derived from human cervical
CC epithelial cells. By measuring gene expression, the probes are therefore
CC useful in grading and/or staging of diseases of the cervix, notably
CC cervical cancer. Note: The sequence data for this patent did not form
CC part of the printed specification, but was obtained in electronic format
CC directly from WIPO at ftp.wipo.int/pub/published_pct_sequences
XX
XX Sequence 48 AA;
XX
XX Query Match          33.0%; Score 32; DB 4; Length 48;
XX Best Local Similarity 33.3%; Pred. No. 6.3e+02;
XX Matches 5; Conservative 4; Mismatches 6; Indels 0; Gaps 0;

QY 3 NHLNSKIAFKIVSQE 17
   ||:|:|:|:|:|:|
Db 14 NHINVMVKFPSIVEE 28

RESULT 30
AAB32791
ID AAM26253 standard; protein; 48 AA.
XX
AC AAM26253;
XX
XX 17-OCT-2001 (first entry)
XX
DE Peptide #290 encoded by probe for measuring placental gene expression.
XX
KW Probe; microarray; human; placenta; antenatal diagnosis;
KW genetic disorder.
XX
OS Homo sapiens.
XX
PN WO200157272-A2.
XX
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XX PD 09-AUG-2001.  
 XX PF 30-JAN-2001; 2001WO-US000663.  
 XX PR 04-FEB-2000; 2000US-0180312P.  
 XX PR 26-MAY-2000; 2000US-0207456P.  
 XX PR 30-JUN-2000; 2000US-00608408.  
 XX PR 03-AUG-2000; 2000US-00632366.  
 XX PR 21-SEP-2000; 2000US-0234687P.  
 XX PR 27-SEP-2000; 2000US-0236359P.  
 XX PR 04-OCT-2000; 2000GB-00024263.  
 XX PA (MOLE-) MOLECULAR DYNAMICS INC.  
 XX PI Penn SG, Hanzel DK, Chen W, Rank DR;  
 XX WPI; 2001-488897/53.  
 XX PT Human genome-derived single exon nucleic acid probes useful for analyzing  
 XX PT gene expression in human placenta.  
 XX PS Claim 27; SEQ ID NO 26522; 654pp; English.  
 XX CC The present invention relates to single exon nucleic acid probes (SENP;  
 CC see AA131315-AA157546). The present sequence is a peptide encoded by one  
 CC such probe. The probes are useful for producing a microarray for  
 CC predicting, measuring and displaying gene expression in samples derived  
 CC from human placenta. The probes are useful for antenatal diagnosis of  
 CC human genetic disorders  
 XX SQ Sequence 48 AA;  
 Query Match 33.0%; Score 32; DB 4; Length 48;  
 Best Local Similarity 33.3%; Pred. No. 6.3e+02;  
 Matches 5; Conservative 4; Mismatches 6; Indels 0; Gaps 0;  
 QY 3 NHLSKIAFKIVSQE 17  
 ||:|:|:|:|:|:|:  
 Db 14 NHINVMVKFPSIVEE 28  
 RESULT 32  
 ABS27621  
 ID ABB27621 standard; peptide; 48 AA.  
 XX AC ABB27621;  
 XX DT 01-FEB-2002 (first entry)  
 XX DE Human peptide #272 encoded by breast cell single exon nucleic acid probe.  
 XX KW Human; microarray; single exon probe; gene expression; breast; disease;  
 XX KW cancer.  
 XX OS Homo sapiens.  
 XX PN WO200157271-A2.  
 XX PD 09-AUG-2001.  
 XX PF 30-JAN-2001; 2001WO-US000662.  
 XX PR 04-FEB-2000; 2000US-0180312P.  
 XX PR 26-MAY-2000; 2000US-0207456P.  
 XX PR 30-JUN-2000; 2000US-00608408.  
 XX PR 03-AUG-2000; 2000US-00632366.  
 XX PR 21-SEP-2000; 2000US-0234687P.  
 XX PR 27-SEP-2000; 2000US-0236359P.  
 XX PR 04-OCT-2000; 2000GB-00024263.  
 XX PA (MOLE-) MOLECULAR DYNAMICS INC.  
 XX PI Penn SG, Hanzel DK, Chen W, Rank DR;

PI Penn SG, Hanzel DK, Chen W, Rank DR;  
 XX WPI; 2001-496933/54.  
 XX PT New spatially-addressable set of single exon nucleic acid probes, useful  
 XX PT for measuring gene expression in sample derived from human breast,  
 XX PT comprises number of single exon nucleic acid probes.  
 XX PS Claim 27; SEQ ID NO 10589; 327pp + Sequence Listing; English.  
 XX CC The invention relates to a spatially-addressable set of single exon  
 CC nucleic acid probes for measuring gene expression in a sample derived  
 CC from human breast and BT 474 cells. The method involves contacting the  
 CC probes with a collection of detectably labelled nucleic acids derived  
 CC from mRNA of human breast, and then measuring the label bound to each  
 CC probe of the microarray. The probes are useful for verifying the  
 CC expression of regions of genomic DNA predicted to encode proteins. They  
 CC are useful for gene discovery, and for determining predisposition and/or  
 CC prognosing breast disease. Gene expression analysis is useful for  
 CC assessing the toxicity of chemical agents on cells. The microarray of  
 CC this invention presents a far greater diversity of probes for measuring  
 CC gene expression, with far less bias than expressed sequence tag  
 CC microarrays. The method is suitable for rapid production of functional  
 CC information from genomic sequence. The present sequence is a peptide  
 CC encoded by a single exon nucleic acid probe of the invention. Note: The  
 CC sequence data for this patent did not form part of the printed  
 CC specification, but was obtained in electronic format directly from WIPO  
 CC at ftp.wipo.int/pub/published\_pct\_sequences  
 XX SQ Sequence 48 AA;  
 Query Match 33.0%; Score 32; DB 4; Length 48;  
 Best Local Similarity 33.3%; Pred. No. 6.3e+02;  
 Matches 5; Conservative 4; Mismatches 6; Indels 0; Gaps 0;  
 QY 3 NHLSKIAFKIVSQE 17  
 ||:|:|:|:|:|:|:  
 Db 14 NHINVMVKFPSIVEE 28  
 RESULT 33  
 ABB18273  
 ID ABB18273 standard; protein; 48 AA.  
 XX AC ABB18273;  
 XX DT 23-JAN-2002 (first entry)  
 XX DE Protein #272 encoded by probe for measuring heart cell gene expression.  
 XX KW Human; gene expression; heart; microarray; vascular system;  
 XX KW cardiovascular disease; hypertension; cardiac arrhythmia;  
 XX KW congenital heart disease.  
 XX OS Homo sapiens.  
 XX PN WO200157274-A2.  
 XX PD 09-AUG-2001.  
 XX PF 30-JAN-2001; 2001WO-US000666.  
 XX PR 04-FEB-2000; 2000US-0180312P.  
 XX PR 26-MAY-2000; 2000US-0207456P.  
 XX PR 30-JUN-2000; 2000US-00608408.  
 XX PR 03-AUG-2000; 2000US-00632366.  
 XX PR 21-SEP-2000; 2000US-0234687P.  
 XX PR 27-SEP-2000; 2000US-0236359P.  
 XX PR 04-OCT-2000; 2000GB-00024263.  
 XX PA (MOLE-) MOLECULAR DYNAMICS INC.  
 XX PI Penn SG, Hanzel DK, Chen W, Rank DR;

XX WPI; 2001-488899/53.  
XX Single exon nucleic acid probes for analyzing gene expression in human  
PT hearts.  
XX  
XX Claim 15; SEQ ID NO 20043; 530pp; English.  
XX  
XX The present invention relates to single exon nucleic acid probes for  
CC measuring human gene expression in a sample derived from human heart (see  
CC ABA21535-ABAA1305). The present sequence is a protein encoded by one such  
CC probe. The probes may be used for predicting, measuring and displaying  
CC gene expression in samples derived from the human heart via microarrays.  
CC By measuring gene expression, the probes are useful for predicting,  
CC diagnosing, grading, staging, monitoring and prognosing diseases of the  
CC human heart and vascular system e.g. cardiovascular disease,  
CC hypertension, cardiac arrhythmias and congenital heart disease. Note: The  
CC sequence data for this patent did not form part of the printed  
CC specification, but was obtained in electronic format directly from WIPO  
CC at ftp.wipo.int/pub/published\_pct\_sequences  
XX  
SQ Sequence 48 AA;  
Query Match 33.0%; Score 32; DB 4; Length 48;  
Best Local Similarity 33.3%; Pred. No. 6.3e+02;  
Matches 5; Conservative 4; Mismatches 6; Indels 0; Gaps 0;  
Qy 3 NHLNSKIAFKIVSQE 17  
Db 14 NHINVMVKFPIVSEE 28  
RESULT 35  
AAM53598  
ID AAM53598 standard; protein; 48 AA.  
XX  
XX AAM53598;  
XX  
XX 05-NOV-2001 (first entry)  
XX Human brain expressed single exon probe encoded protein SEQ ID NO: 25703.  
XX Human; brain expressed exon; gene expression analysis; probe; microarray;  
XX Alzheimer's disease; multiple sclerosis; schizophrenia; epilepsy; cancer.  
XX Homo sapiens.  
XX WO200157275-A2.  
XX  
XX 09-AUG-2001.  
XX  
XX 30-JAN-2001; 2001WO-US0000667.  
XX  
XX 04-FEB-2000; 2000US-0180312P.  
XX 26-MAY-2000; 2000US-0207456P.  
XX 30-JUN-2000; 2000US-00608408.  
XX 03-AUG-2000; 2000US-00632366.  
XX 21-SEP-2000; 2000US-0234687P.  
XX 27-SEP-2000; 2000US-0236359P.  
XX 04-OCT-2000; 2000GB-00024263.  
XX (MOLE-) MOLECULAR DYNAMICS INC.  
XX Penn SG, Hanzel DK, Chen W, Rank DR;  
XX WPI; 2001-483446/52.  
XX  
XX Single exon nucleic acid probes for analyzing gene expression in human  
PT brains.  
XX  
XX Example 4; SEQ ID NO 25703; 650pp + Sequence Listing; English.  
XX  
XX The present invention provides a number of single exon nucleic acid  
CC probes which are derived from genomic sequences expressed in the human  
CC brain. They can be used to measure gene expression in brain cell samples,  
CC which may enable the diagnosis and improved treatment of nervous system  
CC diseases such as Alzheimer's disease, multiple sclerosis, schizophrenia, f  
CC epilepsy and cancers. The present sequence is a protein encoded by one of  
CC the probes of the invention  
XX  
SQ Sequence 48 AA;  
Query Match 33.0%; Score 32; DB 4; Length 48;  
Best Local Similarity 33.3%; Pred. No. 6.3e+02;  
Matches 5; Conservative 4; Mismatches 6; Indels 0; Gaps 0;  
Qy 3 NHLNSKIAFKIVSQE 17  
Db 14 NHINVMVKFPIVSEE 28

XX WPI; 2001-488899/53.  
XX Single exon nucleic acid probes for analyzing gene expression in human  
PT hearts.  
XX  
XX Claim 15; SEQ ID NO 20043; 530pp; English.  
XX  
XX The present invention relates to single exon nucleic acid probes for  
CC measuring human gene expression in a sample derived from human heart (see  
CC ABA21535-ABAA1305). The present sequence is a protein encoded by one such  
CC probe. The probes may be used for predicting, measuring and displaying  
CC gene expression in samples derived from the human heart via microarrays.  
CC By measuring gene expression, the probes are useful for predicting,  
CC diagnosing, grading, staging, monitoring and prognosing diseases of the  
CC human heart and vascular system e.g. cardiovascular disease,  
CC hypertension, cardiac arrhythmias and congenital heart disease. Note: The  
CC sequence data for this patent did not form part of the printed  
CC specification, but was obtained in electronic format directly from WIPO  
CC at ftp.wipo.int/pub/published\_pct\_sequences  
XX  
SQ Sequence 48 AA;  
Query Match 33.0%; Score 32; DB 4; Length 48;  
Best Local Similarity 33.3%; Pred. No. 6.3e+02;  
Matches 5; Conservative 4; Mismatches 6; Indels 0; Gaps 0;  
Qy 3 NHLNSKIAFKIVSQE 17  
Db 14 NHINVMVKFPIVSEE 28  
RESULT 34  
AAM65977  
ID AAM65977 standard; protein; 48 AA.  
XX  
XX AAM65977;  
XX  
XX 06-NOV-2001 (first entry)  
XX Human bone marrow expressed probe encoded protein SEQ ID NO: 26283.  
XX Human; bone marrow expressed exon; gene expression analysis; probe;  
XX microarray; cancer; leukaemia; lymphoma; myeloma.  
XX Homo sapiens.  
XX WO200157276-A2.  
XX  
XX 09-AUG-2001.  
XX  
XX 30-JAN-2001; 2001WO-US0000668.  
XX  
XX 04-FEB-2000; 2000US-0180312P.  
XX 26-MAY-2000; 2000US-0207456P.  
XX 30-JUN-2000; 2000US-00608408.  
XX 03-AUG-2000; 2000US-00632366.  
XX 21-SEP-2000; 2000US-0234687P.  
XX 27-SEP-2000; 2000US-0236359P.  
XX 04-OCT-2000; 2000GB-00024263.  
XX (MOLE-) MOLECULAR DYNAMICS INC.  
XX Penn SG, Hanzel DK, Chen W, Rank DR;  
XX WPI; 2001-488900/53.  
XX  
XX Human genome-derived single exon nucleic acid probes useful for analyzing  
PT gene expression in human bone marrow.  
XX  
XX Example 4; SEQ ID NO 26283; 650pp + Sequence Listing; English.  
XX  
XX The present invention provides a number of single exon nucleic acid

```

Db      14 NHINVMVKFPSIVEE 28

RESULT 36
ABG47643
ID   ABG47643 standard; peptide; 48 AA.
AC
XX
AC   ABG47643;
XX
DT   25-FEB-2003 (first entry)
XX
DE   Human liver peptide, SEQ ID NO 26291.
XX
DE   Human; liver; cirrhosis; hyperlipoproteinaemia; hyperlipidaemia;
XX   hypercholesterolaemia; coronary heart disease.
XX
OS   Homo sapiens.
XX
XX
XX   WO200157273-A2.
XX
XX   09-AUG-2001.
XX
XX   30-JAN-2001; 2001WO-US000664.
XX
XX   04-FEB-2000; 2000US-0180312P.
XX   26-MAY-2000; 2000US-0207456P.
XX   30-JUN-2000; 2000US-00608408.
XX   03-AUG-2000; 2000US-00632366.
XX   21-SEP-2000; 2000US-0234687P.
XX   27-SEP-2000; 2000US-0236359P.
XX   04-OCT-2000; 2000GB-00024263.
XX
XX   (MOLE-) MOLECULAR DYNAMICS INC.
XX
XX   Penn SG, Hanzel DK, Chen W, Rank DR;
XX   WPI; 2001-488898/53.
XX
XX   Human genome-derived single exon nucleic acid probes useful for analyzing
XX   gene expression in human adult liver.
XX
XX   Claim 27; SEQ ID NO 26291; 658pp; English.
XX
XX   The invention relates to a single exon nucleic acid probe (SENP) (I) for
XX   measuring human gene expression in a sample derived from human adult
XX   liver, comprising one of 13109 defined nucleotide sequences given in the
XX   specification (or complements/ fragments). The probe hybridises at high
XX   stringency to a nucleic acid molecule expressed in the human adult liver.
XX   (I) may be used for predicting, measuring and displaying gene expression
XX   in samples derived from human adult liver. The genes identified may be
XX   involved in genetic liver diseases such as cirrhosis,
XX   hyperlipoproteinaemia, hyperlipidaemia and hypercholesterolaemia which is
XX   associated with coronary heart disease. ABG47348-ABG5930 represent human
XX   liver single exon encoded peptides of the invention. Note: The sequence
XX   information for this patent does not appear in the printed specification
XX   but was obtained in electronic format directly from WIPO at
XX   ftp.wipo.int/pub/published_pct_sequences
XX
XX   Sequence 48 AA;
XX
XX   Query Match      33.0%; Score 32; DB 4; Length 48;
XX   Best Local Similarity 33.3%; Pred. No. 6.3e+02;
XX   Matches 5; Conservative 4; Mismatches 6; Indels 0; Gaps 0;

QY      3 NHLNSKIAFKIVSQE 17
Db      14 NHINVMVKFPSIVEE 28

RESULT 37
AAW01589
ID   AAW01589 standard; protein; 48 AA.
XX
XX
XX   Peptide #271 encoded by probe for measuring human breast gene expression.
XX
XX   Probe; human; breast disease; breast cancer; development disorder;
XX   inflammatory disease; proliferative breast disease; non-carcinoma tumour.
XX
XX   Homo sapiens.
XX
XX   WO200157270-A2.
XX
XX   09-AUG-2001.
XX
XX   29-JAN-2001; 2001WO-US000661.
XX
XX   04-FEB-2000; 2000US-0180312P.
XX   26-MAY-2000; 2000US-0207456P.
XX   30-JUN-2000; 2000US-00608408.
XX   03-AUG-2000; 2000US-00632366.
XX   21-SEP-2000; 2000US-0234687P.
XX   27-SEP-2000; 2000US-0236359P.
XX   04-OCT-2000; 2000GB-00024263.
XX
XX   (MOLE-) MOLECULAR DYNAMICS INC.
XX
XX   Penn SG, Hanzel DK, Chen W, Rank DR;
XX   WPI; 2001-476286/51.
XX
XX   Novel single exon nucleic acid probe used to measuring gene expression in
XX   a human breast.
XX
XX   Claim 27; SEQ ID NO 10329; 322pp; English.
XX
XX   The present invention relates to novel single exon nucleic acid probes
XX   (see AA00010-A110067). The present sequence is a peptide encoded by one
XX   a human probe. The probes are useful for measuring human gene expression in
XX   a human breast sample, where the probe hybridises at high stringency to a
XX   nucleic acid expressed in the human breast. The probes are useful for
XX   predicting, diagnosing, grading, staging, monitoring and prognosing
XX   diseases of the human breast, particularly those diseases with polygenic
XX   aetiology. The diseases include: breast cancer, disorders of development,
XX   inflammatory diseases of the breast, fibrocystic changes, proliferative,
XX   breast disease and non-carcinoma tumours. Note: The sequence data for
XX   this patent did not form part of the printed specification, but was
XX   obtained in electronic format directly from WIPO at
XX   ftp.wipo.int/pub/published_pct_sequences
XX
XX   Sequence 48 AA;
XX
XX   Query Match      33.0%; Score 32; DB 4; Length 48;
XX   Best Local Similarity 33.3%; Pred. No. 6.3e+02;
XX   Matches 5; Conservative 4; Mismatches 6; Indels 0; Gaps 0;

QY      3 NHLNSKIAFKIVSQE 17
Db      14 NHINVMVKFPSIVEE 28

RESULT 38
ABG35625
ID   ABG35625 standard; peptide; 48 AA.
XX
XX
XX   AC   ABG35625;
XX
XX   DT   19-AUG-2002 (first entry)
XX
XX   Human peptide encoded by genome-derived single exon probe SEQ ID 25290.
XX
XX   Human; single exon probe; asthma; lung cancer; COPD; ILD;
XX   chronic obstructive pulmonary disease; interstitial lung disease;

```



```

XX SQ Sequence 48 AA;
Query Match          33.0%; Score 32; DB 5; Length 48;
Best Local Similarity 33.3%; Pred. NO. 6.3e+02;
Matches 5; Conservative 4; Mismatches 6; Indels 0; Gaps 0;

QY      3 NHIKSKIAPKIVSQE 17
       ||| : | : | i : |
Db      14 NHINVMVKPPSIVEE 28

RESULT 39
AAB89226
ID AAB89226 standard; peptide; 21 AA.
XX AC AAB89226;
DT DT 23-MAY-2001 (first entry)
XX XX
DE DE HIV gp120 protein binding peptide #319.
XX KW Human chemokine receptor; CD4; HIV; glycoprotein 120; gp120; antagonist;
XX KW replication; CCR5; CXCR4; CD4; STRL33.
XX OS Homo sapiens.
XX PN WO200116182-A2.
XX PD 08-MAR-2001.
XX XX
PF PF 25-AUG-2000; 2000WO-US023505.
XX PR 27-AUG-1999; 99US-0151270P.
XX PA (USSH ) US DEPT HEALTH & HUMAN SERVICES.
XX PI Saxinger C;
XX XX
DR DR WPI; 2001-244398/25.
XX XX
PT PT Novel polypeptides useful for treating HIV infection, have homology to
PT PT regions of domains of human chemokine receptors CCR5, CXCR4 and STRL33,
PT PT and binds to HIV gp120 under physiological conditions.
XX XX
PF PF Example 4; Page 46; 114pp; English.
XX PS The present invention describes a number of peptides which are able to
XX CC bind to HIV glycoprotein 120 (gp120). These are similar to the human
XX CC chemokine receptors CCR5, CXCR4 and STRL33, as well as CD4. These are
XX CC useful in the treatment of HIV, as they prevent replication of the virus.
XX CC The present sequence is an example of a peptide of the invention
XX SQ Sequence 21 AA;

Query Match          32.0%; Score 31; DB 4; Length 21;
Best Local Similarity 38.5%; Pred. NO. 3.5e+02;
Matches 5; Conservative 5; Mismatches 3; Indels 0; Gaps 0;

QY      6 NSKIAPKIVSQEP 18
       | : : | | : |
Db      8 NKEVSVKRVTQDP 20

RESULT 40
AAB89227
ID AAB89227 standard; peptide; 21 AA.
XX AC AAB89227;
XX XX
DT DT 23-MAY-2001 (first entry)
XX XX
DE DE HIV gp120 protein binding peptide #320.

```

XX Human chemokine receptor; CD4; HIV; glycoprotein 120; gp120; antagonist;  
 KW replication; CCR5; CXCR4; CD4; STRL33.  
 XX Homo sapiens.  
 OS  
 XX WO200116182-A2.  
 FN  
 XX 08-MAR-2001.  
 PD  
 XX 25-AUG-2000; 2000WO-US023505.  
 PF  
 XX 27-AUG-1999; 99US-0151270P.  
 PR  
 XX (USSH ) US DEPT HEALTH & HUMAN SERVICES.  
 XX Saxinger C;  
 XX WPI; 2001-244398/25.  
 DR Novel polypeptides useful for treating HIV infection, have homology to  
 PT regions of domains of human chemokine receptors CCR5, CXCR4 and STRL33,  
 PT and binds to HIV gp120 under physiological conditions.  
 XX Example 4; Page 46; 114pp; English.  
 PS  
 XX The present invention describes a number of peptides which are able to  
 CC bind to HIV glycoprotein 120 (gp120). These are similar to the human  
 CC chemokine receptors CCR5, CXCR4 and STRL33, as well as CD4. These are  
 CC useful in the treatment of HIV, as they prevent replication of the virus.  
 CC The present sequence is an example of a peptide of the invention  
 XX  
 SQ Sequence 21 AA;

Query Match 32.0%; Score 31; DB 4; Length 21;  
 Best Local Similarity 38.5%; Pred. No. 3.5e+02;  
 Matches 5; Conservative 5; Mismatches 3; Indels 0; Gaps 0;  
 QY 6 NSKIAFKIVSQEP 18  
 | : : : | : : : |  
 Db 3 NKEVSVKRVITQDP 15

RESULT 41  
 AAY27522  
 ID AAY27522 standard; peptide; 24 AA.  
 XX  
 AC AAY27522;  
 XX  
 DT 29-NOV-1999 (first entry)  
 XX  
 DE E. coli beta'-subunit conserved regions B and I derived peptide 55.  
 XX  
 KW Bacterial life cycle; protein subunit; ribonucleic acid polymerase; RNAP;  
 KW enzyme function; anti-bacterial; E. coli.  
 XX  
 OS Synthetic.  
 OS Escherichia coli.  
 XX  
 FN WO9943338-A1.  
 XX  
 PD 02-SEP-1999.  
 XX  
 PF 26-FEB-1999; 99WO-US0004351.  
 XX  
 PR 26-FEB-1998; 98US-00031122.  
 XX  
 PA (GLIN/) GLINSKII G V.  
 XX  
 PI Glinskii GV;  
 XX  
 DR WPI; 1999-550829/46.  
 XX

PT Identifying antibacterial drugs, by identifying compounds that block the  
 PT binding of protein subunits of ribonucleic acid polymerase.  
 XX Claim 31; Page 64; 106pp; English.

XX The invention relates to methods of interfering with bacterial life cycle  
 CC by bringing bacterial cells into contact with a compound that blocks the  
 CC binding of at least one protein subunit of ribonucleic acid polymerase  
 CC (RNAP) to a second protein subunit of RNAP. The methods can be used for  
 CC obtaining compounds which inhibit subunit-subunit interactions and  
 CC assembly necessary for enzyme function in bacteria. The compounds inhibit  
 CC the binding of (a) at least one protein subunit of RNAP to a second  
 CC protein subunit of RNAP; (b) the sigma-subunit of RNAP to the RNAP core;  
 CC and blocks a nucleic acid binding to the beta-subunit or the beta'-  
 CC subunit of RNAP. The compounds obtained can be used as anti-bacterial  
 CC drugs. Sequences AAY27509-523 represent peptides derived from the  
 CC putative nucleic acid binding sequences of the conserved regions B and I  
 CC of E. coli beta'-subunit. The antibacterial compounds that block nucleic  
 CC acid binding to the beta-subunit of the RNAP bind to the sequences  
 CC indicated above  
 XX  
 SQ Sequence 24 AA;

Query Match 32.0%; Score 31; DB 2; Length 24;  
 Best Local Similarity 33.3%; Pred. No. 4.1e+02;  
 Matches 5; Conservative 5; Mismatches 5; Indels 0; Gaps 0;  
 QY 4 HLNSKIAFKIVSQEP 18  
 | : : : | : : : |  
 Db 2 HARSTGYSILVTQDP 16

RESULT 42  
 AAY02263  
 ID AAY02263 standard; protein; 33 AA.  
 XX  
 AC AAY02263;  
 XX  
 DT 08-JUL-1999 (first entry)  
 XX  
 DE A F-box protein sequence.  
 XX  
 KW F-box protein; targeted ubiquitination; cellular protein;  
 KW cell cycle regulator; transcription regulator; DNA replication;  
 KW inflammatory response; infectious disease; protein degradation; cancer;  
 KW virus infection.  
 XX  
 OS Mus sp.  
 XX  
 PN WO9918989-A1.  
 XX  
 PD 22-APR-1999.  
 XX  
 PF 15-OCT-1998; 98WO-US021763.  
 XX  
 PR 16-OCT-1997; 97US-00951621.  
 XX  
 PA (BAYU ) BAYLOR COLLEGE MEDICINE.  
 XX  
 PI Harper JW, Ellledge SJ;  
 XX  
 DR WPI; 1999-277441/23.  
 DR N-PSDB; AAX35537.  
 XX  
 PT New isolated F-box proteins and genes for development of therapeutics,  
 PT e.g. for cancer treatment.  
 XX  
 PS Claim 4; Page 109; 170pp; English.

XX AAX35523-51 encode F-box proteins (AAY02249-77) which are involved in the  
 CC targeted ubiquitination of cellular proteins. The F-box proteins are  
 CC involved in targeted ubiquitination of cellular proteins, including cell  
 CC cycle regulators. The products and methods can be used for determining

CC the interaction of these proteins with other proteins, e.g. to identify  
 CC and/or investigate cell cycle regulators, transcription regulators,  
 CC proteins involved in DNA replication, and other cellular regulatory  
 CC proteins. They can be used in elucidating inflammatory response and  
 CC infectious disease processes involving protein degradation as well as  
 CC development of compounds that control (i.e. either enhance or retard)  
 CC protein degradation, as appropriate to ameliorate the effects of the  
 CC inflammatory response or disease process. They can be used for  
 CC identifying and developing compounds effective against cancers or virus  
 CC infection, e.g. immunodeficiency viruses such as HIV, feline  
 CC immunodeficiency virus, bovine immunodeficiency virus, and simian  
 CC immunodeficiency virus  
 XX  
 SQ Sequence 33 AA;

Query Match 32.0%; Score 31; DB 2; Length 33;  
 Best Local Similarity 58.3%; Pred. No. 6e+02;  
 Matches 7; Conservative 2; Mismatches 3; Indels 0; Gaps 0;

QY 5 LNSKIAPKIVSQ 16  
 | : | | | | |  
 Db 1 LPAEITFKIFSQ 12

RESULT 43  
 AAE08035 standard; peptide; 33 AA.

AC AAE08035;  
 DT 01-NOV-2001 (first entry)

DE Mouse F-box protein, F10 Lambda.

KW Mouse; nuclear factor-kappaB; NF-kB; regulatory factor; slimb protein;  
 XX targetted ubiquitination; F-box protein; F10 Lambda.

OS Mus musculus.

FN USG232081-B1.

PD 15-MAY-2001.

PF 15-OCT-1998; 98US-00172841.

PR 16-OCT-1997; 97US-00951621.

PA (BAYU ) BAYLOR COLLEGE MEDICINE.

PI Harper JW, Elledge SJ, Winston JT;

DR WPI; 2001-342771/36.

DR N-PSDB; AAD14872.

XX Detecting nuclear factor-kappaB regulatory factors, such as F-box  
 PT proteins involved in targeted ubiquitination, by contacting the  
 PT regulatory factors with slimb protein to form a complex and detecting the  
 PT complex.

PS Example 6; Fig 7; 69pp; English.

XX The present invention relates to a method for detection of one or more  
 CC nuclear factor (NF)-kappaB (kB) regulatory factors. The method comprises  
 CC exposing a slimb protein to a sample suspected of containing one or more  
 CC NF-kB regulatory factors, so that the slimb protein binds to one or more  
 CC NF-kB regulatory factors to form a slimb/regulatory factor complex and  
 CC detecting the slimb/regulatory factor complex. The method is useful for  
 CC detecting NF-kB regulatory factors such as F-box proteins, IxBs, IKKs and  
 CC agonists, antagonists and cofactors that interact with these factors. F-  
 CC box proteins are involved in targetted ubiquitination of cellular  
 CC proteins. The present sequence is mouse F-box protein, F10 Lambda

XX Sequence 33 AA;

Query Match 32.0%; Score 31; DB 4; Length 33;  
 Best Local Similarity 58.3%; Pred. No. 6e+02;  
 Matches 7; Conservative 2; Mismatches 3; Indels 0; Gaps 0;

QY 5 LNSKIAPKIVSQ 16  
 | : | | | | |  
 Db 1 LPAEITFKIFSQ 12

RESULT 44  
 AAE39643 standard; peptide; 33 AA.

AC AAE39643;

DT 18-DEC-2003 (first entry)

DE Mouse F-box protein, F10 (lambda).

KW Mouse; F-Box domain; E3 complex; ubiquitination; cell cycle regulator;  
 XX inflammatory disease.

OS Mus sp.

FN US6573094-B1.

PD 03-JUN-2003.

PF 16-OCT-1997; 97US-00951621.

PR 16-OCT-1997; 97US-00951621.

PA (BAYU ) BAYLOR COLLEGE MEDICINE.

PI Harper JW, Elledge SJ;

DR WPI; 2003-776006/73.

DR N-PSDB; AAD60312.

XX New isolated nucleic acid segment encoding a protein with at least one  
 PT functionally active F-box domain, useful for identifying related genes,  
 PT and for developing compounds for treating infectious or inflammatory  
 PT disease.

PS Example 6; Col 57-58; Opp; English.

XX The invention relates to an isolated nucleic acid segment comprising or  
 CC consisting essentially of a nucleic acid sequence encoding a protein  
 CC comprising at least one functionally active F-Box domain sequence. The  
 CC polypeptide encoded by the nucleic acid segment is part of an E3 complex  
 CC involved in ubiquitination of cell cycle regulators and may be useful in  
 CC investigating mechanisms of infectious and inflammatory diseases and in  
 CC developing therapeutic agents for treating such diseases. The invention  
 CC is useful for detecting related polynucleotides encoding F-box proteins  
 CC and in the determination of the function of proteins such as elongin C,  
 CC Skp1-related protein, elongin B and elongin A. The present sequence is  
 CC mouse F-box protein

XX Sequence 33 AA;

Query Match 32.0%; Score 31; DB 7; Length 33;  
 Best Local Similarity 58.3%; Pred. No. 6e+02;  
 Matches 7; Conservative 2; Mismatches 3; Indels 0; Gaps 0;

QY 5 LNSKIAPKIVSQ 16  
 | : | | | | |  
 Db 1 LPAEITFKIFSQ 12

RESULT 45  
 AAW73422  
 ID AAW73422 standard; protein; 34 AA.  
 XX AC AAW73422;  
 XX 19-FEB-1999 (first entry)  
 DT  
 DE Human secreted protein encoded by Gene No. 26.  
 XX  
 XX Secreted protein; human; protein therapy; gene therapy; blood disorder;  
 KW pathological condition; diagnosis; cancer; neurological disorder;  
 KW developmental abnormality; foetal deficiency; leukaemia; hepatic disease;  
 KW immune system disorder; Alzheimer's disease; cognitive disorder;  
 KW schizophrenia; prostate disease; autoimmune disease; AIDS.  
 XX  
 OS Homo sapiens.  
 XX  
 XX Key Location/Qualifiers  
 FH Misc-difference 34  
 FT /note= "unspecified amino acid"  
 XX  
 XX WO9854206-A1.  
 XX  
 XX 03-DEC-1998.  
 XX  
 XX 28-MAY-1998; 98WO-US010868.  
 XX  
 XX 30-MAY-1997; 97US-0044039P.  
 XX 30-MAY-1997; 97US-0048093P.  
 XX 30-MAY-1997; 97US-0048101P.  
 XX 30-MAY-1997; 97US-0048190P.  
 XX 30-MAY-1997; 97US-0048356P.  
 XX 30-MAY-1997; 97US-0050935P.  
 XX 29-AUG-1997; 97US-0056250P.  
 XX 29-AUG-1997; 97US-0056293P.  
 XX 29-AUG-1997; 97US-0056296P.  
 XX  
 XX (HUMA-) HUMAN GENOME SCI INC.  
 XX  
 XX Ruben SM, Rosen CA, Carter KC, Dillon PJ, Endress GA, Yu G;  
 PI Ni J, Feng P;  
 XX  
 XX WPI: 1999-070209/06.  
 XX N-PSDB; AAV08836.  
 DR  
 DR  
 XX  
 XX New isolated human genes - useful for diagnosis and treatment of, e.g.  
 PT cancers, neurological disorders, immune diseases, developmental disorders  
 PT or blood disorders.  
 PT  
 PS Claim 11; Page 160; 198pp; English.  
 XX  
 CC This sequence is encoded by a cDNA of the invention, designated Gene No.  
 CC 26. This sequence represents a human secreted protein, and is expressed  
 CC in a variety of tissues including colon cancer, breast cancer,  
 CC neutrophils, T-cells, spinal fluid, fibroblasts and vascular endothelial  
 CC cells. The DNA sequences of the invention and their corresponding  
 CC secreted polypeptides are useful for preventing, treating or ameliorating  
 CC medical conditions, e.g. by protein or gene therapy. Also pathological  
 CC conditions can be diagnosed by determining the amount of the new  
 CC polypeptides in a sample or by determining the presence of mutations in  
 CC the DNA sequences. Specific uses are described for each of the DNA  
 CC sequences and the encoded proteins, based on which tissues they are most  
 CC highly expressed in, and include developing products for the diagnosis or  
 CC treatment of cancer, tumours, neurological disorders, developmental  
 CC abnormalities and foetal deficiencies, blood disorders, leukaemias,  
 CC diseases of the immune system (including allergies or asthma), hepatic  
 CC disease, Alzheimer's and cognitive disorders, schizophrenia, prostate  
 CC diseases, autoimmune disorders and AIDS. The polypeptides are also useful  
 CC for identifying their binding partners  
 XX  
 SQ Sequence 34 AA;  
 XX

Query Match 32.0%; Score 31; DB 2; Length 34;  
 Best Local Similarity 47.4%; Pred. No. 6.2e+02;  
 Matches 9; Conservative 2; Mismatches 4; Indels 1;  
 QY 5 LNSKIAPKIV---SOEPA 19  
 |||||:|:|  
 Db 11 LNSKLVAAVVNLIKASQMPA 29  
 |||||:|:|  
 RESULT 46  
 AAM95698  
 ID AAM95698 standard; protein; 34 AA.  
 XX AC AAM95698;  
 XX 21-NOV-2001 (first entry)  
 DT  
 DE Human reproductive system related antigen SEQ ID NO: 4356.  
 XX  
 XX Human; reproductive system related antigen; reproductive system disorder;  
 KW cancer; gene therapy.  
 XX  
 OS Homo sapiens.  
 XX  
 XX WO200155320-A2.  
 XX  
 XX 02-AUG-2001.  
 XX  
 XX 17-JAN-2001; 2001WO-US001339.  
 XX  
 XX 31-JAN-2000; 2000US-0179065P.  
 PR 04-FEB-2000; 2000US-0180628P.  
 PR 24-FEB-2000; 2000US-0184664P.  
 PR 02-MAR-2000; 2000US-0186350P.  
 PR 16-MAR-2000; 2000US-0189874P.  
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 PR 28-JUN-2000; 2000US-0214886P.  
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 PR 11-JUL-2000; 2000US-0217496P.  
 PR 14-JUL-2000; 2000US-0218290P.  
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 PR 14-AUG-2000; 2000US-0225477P.  
 PR 14-AUG-2000; 2000US-0225757P.  
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 PR 18-AUG-2000; 2000US-0226279P.  
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 PR 22-AUG-2000; 2000US-0226688P.  
 PR 22-AUG-2000; 2000US-0227182P.  
 PR 23-AUG-2000; 2000US-0227009P.  
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 PR 01-SEP-2000; 2000US-0228287P.  
 PR 01-SEP-2000; 2000US-0229343P.  
 PR 01-SEP-2000; 2000US-0229344P.  
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 PR 05-SEP-2000; 2000US-0229509P.  
 PR 05-SEP-2000; 2000US-0229513P.



XX PR 31-JAN-2000; 2000US-0179065P.  
PR 04-FEB-2000; 2000US-0180628P.  
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PR 02-MAR-2000; 2000US-0186350P.  
PR 16-MAR-2000; 2000US-0189874P.  
PR 17-MAR-2000; 2000US-0190076P.  
PR 18-APR-2000; 2000US-0198123P.  
PR 19-MAY-2000; 2000US-0205515P.  
PR 07-JUN-2000; 2000US-0209457P.  
PR 28-JUN-2000; 2000US-0214886P.  
PR 30-JUN-2000; 2000US-0215135P.  
PR 07-JUL-2000; 2000US-0216647P.  
PR 07-JUL-2000; 2000US-0216880P.  
PR 11-JUL-2000; 2000US-0217487P.  
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PR 14-AUG-2000; 2000US-0225268P.  
PR 14-AUG-2000; 2000US-0225270P.  
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PR 14-AUG-2000; 2000US-0225757P.  
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PR 18-AUG-2000; 2000US-0226279P.  
PR 22-AUG-2000; 2000US-0226681P.  
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PR 22-AUG-2000; 2000US-0227182P.  
PR 23-AUG-2000; 2000US-0228009P.  
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PR 01-SEP-2000; 2000US-0229287P.  
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PR 05-SEP-2000; 2000US-0229509P.  
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PR 06-SEP-2000; 2000US-0230437P.  
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PR 08-SEP-2000; 2000US-0231243P.  
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PR 08-SEP-2000; 2000US-0231414P.  
PR 08-SEP-2000; 2000US-0232080P.  
PR 08-SEP-2000; 2000US-0232081P.  
PR 12-SEP-2000; 2000US-0231968P.  
PR 14-SEP-2000; 2000US-0232397P.  
PR 14-SEP-2000; 2000US-0232398P.  
PR 14-SEP-2000; 2000US-0232399P.  
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PR 14-SEP-2000; 2000US-0232401P.  
PR 14-SEP-2000; 2000US-0233063P.  
PR 14-SEP-2000; 2000US-0233064P.  
PR 14-SEP-2000; 2000US-0233065P.  
PR 21-SEP-2000; 2000US-0234223P.  
PR 21-SEP-2000; 2000US-0234274P.  
PR 25-SEP-2000; 2000US-0234597P.  
PR 25-SEP-2000; 2000US-0234998P.  
PR 26-SEP-2000; 2000US-0235484P.  
PR 27-SEP-2000; 2000US-0235834P.  
PR 27-SEP-2000; 2000US-0235836P.  
PR 29-SEP-2000; 2000US-0236327P.  
PR 29-SEP-2000; 2000US-0236367P.  
PR 29-SEP-2000; 2000US-0236368P.  
PR 29-SEP-2000; 2000US-0236369P.  
PR 29-SEP-2000; 2000US-0236370P.

PR 02-OCT-2000; 2000US-0236802P.  
PR 02-OCT-2000; 2000US-0237037P.  
PR 02-OCT-2000; 2000US-0237038P.  
PR 02-OCT-2000; 2000US-0237039P.  
PR 02-OCT-2000; 2000US-0237040P.  
PR 13-OCT-2000; 2000US-0239935P.  
PR 13-OCT-2000; 2000US-0239937P.  
PR 20-OCT-2000; 2000US-0240960P.  
PR 20-OCT-2000; 2000US-0241221P.  
PR 20-OCT-2000; 2000US-0241785P.  
PR 20-OCT-2000; 2000US-0241786P.  
PR 20-OCT-2000; 2000US-0241787P.  
PR 20-OCT-2000; 2000US-0241808P.  
PR 20-OCT-2000; 2000US-0241809P.  
PR 20-OCT-2000; 2000US-0241826P.  
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PR 08-NOV-2000; 2000US-0246474P.  
PR 08-NOV-2000; 2000US-0246475P.  
PR 08-NOV-2000; 2000US-0246476P.  
PR 08-NOV-2000; 2000US-0246477P.  
PR 08-NOV-2000; 2000US-0246478P.  
PR 08-NOV-2000; 2000US-0246523P.  
PR 08-NOV-2000; 2000US-0246524P.  
PR 08-NOV-2000; 2000US-0246525P.  
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PR 08-NOV-2000; 2000US-0246527P.  
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PR 08-NOV-2000; 2000US-0246610P.  
PR 08-NOV-2000; 2000US-0246611P.  
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PR 17-NOV-2000; 2000US-0249207P.  
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PR 17-NOV-2000; 2000US-0249213P.  
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PR 17-NOV-2000; 2000US-0249215P.  
PR 17-NOV-2000; 2000US-0249216P.  
PR 17-NOV-2000; 2000US-0249217P.  
PR 17-NOV-2000; 2000US-0249218P.  
PR 17-NOV-2000; 2000US-0249244P.  
PR 17-NOV-2000; 2000US-0249245P.  
PR 17-NOV-2000; 2000US-0249264P.  
PR 17-NOV-2000; 2000US-0249265P.  
PR 17-NOV-2000; 2000US-0249279P.  
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PR 17-NOV-2000; 2000US-0249300P.  
PR 01-DEC-2000; 2000US-0250160P.  
PR 01-DEC-2000; 2000US-0250391P.  
PR 05-DEC-2000; 2000US-0251030P.  
PR 05-DEC-2000; 2000US-0251988P.  
PR 05-DEC-2000; 2000US-0251989P.  
PR 06-DEC-2000; 2000US-0256719P.  
PR 08-DEC-2000; 2000US-0251856P.  
PR 08-DEC-2000; 2000US-0251868P.  
PR 08-DEC-2000; 2000US-0251869P.  
PR 08-DEC-2000; 2000US-0251989P.  
PR 08-DEC-2000; 2000US-0251990P.  
PR 11-DEC-2000; 2000US-0254097P.  
PR 05-JAN-2001; 2001US-0259678P.

(HUMA-) HUMAN GENOME SCI INC.

Rosen CA, Barash SC, Ruben SM;

WPI; 2001-488786/53.

N-PSDB; ABA07588.

PT New isolated ovarian and/or breast cancer related nucleic acids and





CC The invention relates to a nucleic acid probe for measuring human gene  
 CC expression, comprising any of the 27,400 fully defined nucleotide  
 CC sequences in the specification, or their complements or fragments, and  
 CC encoding at least 8 amino acids of any of the 6888 amino acid sequences  
 CC fully defined in the specification. The probe is a single exon probe that  
 CC hybridises under high stringency conditions to a nucleic acid molecule  
 CC expressed in human cells or tissues. Also included are a spatially-  
 CC addressable set of single exon nucleic acid probes for measuring human  
 CC gene expression (comprising a plurality of single exon nucleic acid  
 CC probes cited above, where each of the plurality of probes is separately  
 CC and addressably isolatable or amplifiable from the plurality), a single  
 CC exon microarray for measuring human gene expression, a method of  
 CC measuring human gene expression, a vector comprising the single exon  
 CC probe cited above, an ORF-encoded peptide comprising at least 8  
 CC contiguous amino acids of any of the above-mentioned amino acid  
 CC sequences (optionally with conservative amino acid substitutions), an  
 CC isolated antibody that binds specifically to a peptide cited above,  
 CC a customer desiring to measure gene expression, a method of providing  
 CC human gene expression data by subscription, and a computer-readable  
 CC storage medium which contains a database having a plurality of records  
 CC (each record including data on the expression of a single exon probe  
 CC cited above). The probe, methods and apparatus are useful in gene  
 CC expression analysis. The probes may be used as tools for surveying  
 CC tissues to detect the presence of expressed messages that contain their  
 CC specific exon, or in constructing genome-derived single exon microarrays.  
 CC In addition, the probes are used in identifying and characterising  
 CC alternative splicing events, in detecting and characterising gross  
 CC alterations in the genomic locus that includes their exon, in assessing  
 CC smaller genomic alterations, in priming the synthesis of nucleic acids,  
 CC or in expressing the ORF-encoded peptide. The present sequence is a human  
 CC single exon probe protein of the invention. Note: The sequence data for  
 CC this patent did not form part of the printed specification, but was  
 CC obtained in electronic format directly from USPTO at  
 CC seqdata.uspto.gov/sequence.html?DocID=20030194704

SQ Sequence 43 AA;

Query Match 32.0%; Score 31; DB 8; Length 43;  
 Best Local Similarity 41.7%; Pred. No. 8.3e+02;  
 Matches 5; Conservative 4; Mismatches 3; Indels 0; Gaps 0;

QY 2 PNHLSKIAFKI 13  
 DB 15 PIHLHSNVAWTV 26

RESULT 50  
 AAM21342  
 ID AAM21342 standard; protein; 44 AA.  
 AC AAM21342;  
 AC AAM21342;

DT 12-OCT-2001 (first entry)

DE Peptide #7776 encoded by probe for measuring cervical gene expression.  
 XX Probe; human; microarray; gene expression; cervical epithelial cell;  
 KW cervical cancer.

OS Homo sapiens.

XX WO200157278-A2.

PN 09-AUG-2001.

XX 30-JAN-2001; 2001WO-US000670.

XX 04-FEB-2000; 2000US-0180312P.

PR 26-MAY-2000; 2000US-0207456P.

PR 30-JUN-2000; 2000US-00608408.

PR 03-AUG-2000; 2000US-00632366.

PR 21-SEP-2000; 2000US-0234687P.

PR 27-SEP-2000; 2000US-0236359P.  
 PR 04-OCT-2000; 2000GB-00024263.  
 XX (MOLE-) MOLECULAR DYNAMICS INC.  
 PA Penn SG, Hanzel DK, Chen W, Rank DR;  
 PI WPI; 2001-488901/53.  
 XX Human genome-derived single exon nucleic acid probes useful for analyzing  
 XX gene expression in human cervical epithelial cells.  
 PT Claim 27; SEQ ID NO 26168; 487pp; English.

XX The present invention relates to human single exon nucleic acid probes  
 CC (SENPs: see AAL10068-AA128459). The present sequence is a peptide encoded  
 CC by one such probe. The SENPs are derived from human HeLa cells. The SENPs  
 CC can be used to produce a single exon microarray, which can be used for  
 CC measuring human gene expression in a sample derived from human cervical  
 CC epithelial cells. By measuring gene expression, the probes are therefore  
 CC useful in grading and/or staging of diseases of the cervix, notably  
 CC cervical cancer. Note: The sequence data for this patent did not form  
 CC part of the printed specification, but was obtained in electronic format  
 CC directly from WIPO at ftp.wipo.int/pub/published\_pct\_sequences

SQ Sequence 44 AA;

Query Match 32.0%; Score 31; DB 4; Length 44;  
 Best Local Similarity 33.3%; Pred. No. 8.5e+02;  
 Matches 8; Conservative 3; Mismatches 5; Indels 8; Gaps 1;

QY 3 NHIHNSKIA-----FKIVSQEP 18  
 DB 9 DHINLVAGODGSWQFKIXRHTP 32

RESULT 51

ABB43679

ID ABB43679 standard; peptide; 44 AA.

AC ABB43679;

DT 04-FEB-2002 (first entry)

DE Peptide #1185 encoded by human foetal liver single exon probe.

XX Human; foetal liver; gene expression; single exon nucleic acid probe.

XX Homo sapiens.

XX WO200157277-A2.

PN 09-AUG-2001.

XX 30-JAN-2001; 2001WO-US000669.

XX 04-FEB-2000; 2000US-0180312P.

PR 26-MAY-2000; 2000US-0207456P.

PR 30-JUN-2000; 2000US-00608408.

PR 03-AUG-2000; 2000US-00632366.

PR 21-SEP-2000; 2000US-0234687P.

PR 27-SEP-2000; 2000US-0236359P.

PR 04-OCT-2000; 2000GB-00024263.

XX (MOLE-) MOLECULAR DYNAMICS INC.

XX Penn SG, Hanzel DK, Chen W, Rank DR;

XX WPI; 2001-483447/52.

XX Human genome-derived single exon nucleic acid probes useful for analyzing  
 PT gene expression in human fetal liver.



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AAM77419
ID AAM77419 standard; protein; 44 AA.
XX AC
XX AAM77419;
XX DT
XX 06-NOV-2001 (first entry)
XX DE Human bone marrow expressed probe encoded protein SEQ ID NO: 37725.
XX KW Human; bone marrow expressed exon; gene expression analysis; probe;
XX KW microarray; cancer; leukaemia; lymphoma; myeloma.
XX OS Homo sapiens.
XX XX
XX WO200157276-A2.
XX FN
XX XX
XX PD 09-AUG-2001.
XX XX
XX 30-JAN-2001; 2001WO-US000668.
XX XX
XX 04-FEB-2000; 2000US-0180312P.
XX PR 26-MAY-2000; 2000US-0207456P.
XX PR 30-JUN-2000; 2000US-00608408.
XX PR 03-AUG-2000; 2000US-00632366.
XX PR 21-SEP-2000; 2000US-0234687P.
XX PR 27-SEP-2000; 2000US-0236359P.
XX PR 04-OCT-2000; 2000GB-00024263.
XX XX
XX (MOLE-) MOLECULAR DYNAMICS INC.
XX PA
XX Penn SG, Hanzel DK, Chen W, Rank DR;
XX FI
XX WPI; 2001-483446/53.
XX DR
XX XX
XX Single exon nucleic acid probes for analyzing gene expression in human
XX PT brains.
XX PT
XX Example 4; SEQ ID NO 37725; 658pp + Sequence Listing; English.
XX PS
XX CC The present invention provides a number of single exon nucleic acid
XX CC probes which are derived from genomic sequences expressed in the human
XX CC brain. They can be used to measure gene expression in bone marrow
XX CC samples, which may enable the improved diagnosis and treatment of cancers
XX CC such as lymphoma, leukaemia and myeloma. The present sequence is a
XX CC protein encoded by one of the probes of the invention
XX CC
XX SQ Sequence 44 AA;

Query Match 32.0%; Score 31; DB 4; Length 44;
Best Local Similarity 33.3%; Pred. No. 8.5e+02;
Matches 8; Conservative 3; Mismatches 5; Indels 8; Gaps 1;

XX
XX 3 NHLNKSIA-----FKIVSQP 18
Db 9 DHINLVAGDGSVQFKIKRHTP 32

RESULT 55
ID AAM64636 standard; protein; 44 AA.
XX AC
XX AAM64636;
XX DT
XX 05-NOV-2001 (first entry)
XX DE Human brain expressed single exon probe encoded protein SEQ ID NO: 36741.
XX KW Human; brain expressed exon; gene expression analysis; probe; microarray;
XX KW Alzheimer's disease; multiple sclerosis; schizophrenia; cancer.
XX XX
XX OS Homo sapiens.
XX XX
XX WO200157275-A2.
XX FN
XX XX
XX PD 09-AUG-2001.
XX XX
XX 30-JAN-2001; 2001WO-US000664.
XX XX
XX 04-FEB-2000; 2000US-0180312P.
XX PR 26-MAY-2000; 2000US-0207456P.
XX PR 30-JUN-2000; 2000US-00608408.
XX PR 03-AUG-2000; 2000US-00632366.
XX PR 21-SEP-2000; 2000US-0234687P.
XX PR 27-SEP-2000; 2000US-0236359P.
XX PR 04-OCT-2000; 2000GB-00024263.
XX XX
XX (MOLE-) MOLECULAR DYNAMICS INC.
XX PA
XX Penn SG, Hanzel DK, Chen W, Rank DR;
XX FI
XX WPI; 2001-483446/52.
XX DR
XX XX
XX Single exon nucleic acid probes for analyzing gene expression in human
XX PT brains.
XX PT
XX Example 4; SEQ ID NO 36741; 650pp + Sequence Listing; English.
XX PS
XX CC The present invention provides a number of single exon nucleic acid
XX CC probes which are derived from genomic sequences expressed in the human
XX CC brain. They can be used to measure gene expression in brain cell samples,
XX CC which may enable the diagnosis and improved treatment of nervous system
XX CC diseases such as Alzheimer's disease, multiple sclerosis, schizophrenia,
XX CC epilepsy and cancers. The present sequence is a protein encoded by one of
XX CC the probes of the invention
XX CC
XX SQ Sequence 44 AA;

Query Match 32.0%; Score 31; DB 4; Length 44;
Best Local Similarity 33.3%; Pred. No. 8.5e+02;
Matches 8; Conservative 3; Mismatches 5; Indels 8; Gaps 1;

XX
XX 3 NHLNKSIA-----FKIVSQP 18
Db 9 DHINLVAGDGSVQFKIKRHTP 32

RESULT 56
ID ABG59052 standard; peptide; 44 AA.
XX AC
XX ABG59052;
XX DT
XX 25-FEB-2003 (first entry)
XX DE Human liver peptide, SEQ ID NO 37700.
XX KW Human; liver; cirrhosis; hyperlipoproteinaemia; hyperlipidaemia;
XX KW hypercholesterolaemia; coronary heart disease.
XX XX
XX OS Homo sapiens.
XX XX
XX WO200157273-A2.
XX FN
XX XX
XX PD 09-AUG-2001.
XX XX
XX 30-JAN-2001; 2001WO-US000664.
XX XX
XX 04-FEB-2000; 2000US-0180312P.
XX PR 26-MAY-2000; 2000US-0207456P.
XX PR 30-JUN-2000; 2000US-00608408.
XX PR 03-AUG-2000; 2000US-00632366.
XX PR 21-SEP-2000; 2000US-0234687P.
XX PR 27-SEP-2000; 2000US-0236359P.
XX PR 04-OCT-2000; 2000GB-00024263.
XX XX
XX (MOLE-) MOLECULAR DYNAMICS INC.
XX PA

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XX Penn SG, Hanzel DK, Chen W, Rank DR;  
XX WPI; 2001-48898/53.  
XX Human genome-derived single exon nucleic acid probes useful for analyzing  
PT gene expression in human adult liver.  
XX Claim 27; SEQ ID NO 37700; 658pp; English.  
XX The invention relates to a single exon nucleic acid probe (SENp) (I) for  
CC measuring human gene expression in a sample derived from human adult  
CC liver, comprising one of 13109 defined nucleotide sequences given in the  
CC specification (or complements/ fragments). The probe hybridises at high  
CC stringency to a nucleic acid molecule expressed in the human adult liver.  
CC (I) may be used for predicting, measuring and displaying gene expression  
CC in samples derived from human adult liver. The genes identified may be  
CC involved in genetic liver diseases such as cirrhosis,  
CC hyperlipoproteinaemia, hyperlipidaemia and hypercholesterolaemia which is  
CC associated with coronary heart diseases. ABG47348-ABG59930 represent human  
CC liver single exon encoded peptides of the invention. Note: the sequence  
CC information for this patent does not appear in the printed specification  
CC but was obtained in electronic format directly from WIPO at  
CC ftp.wipo.int/pub/published\_pct\_sequences  
XX Sequence 44 AA;  
SQ  
Query Match 32.0%; Score 31; DB 4; Length 44;  
Best Local Similarity 33.3%; Pred. No. 8.5e+02;  
Matches 8; Conservative 3; Mismatches 5; Indels 8; Gaps 1;  
QY 3 NMLNSKIA-----FKIVSQEP 18  
Db 9 DHINLKVAGQGSVVQVFKIKRHTP 32  
RESULT 57  
AAU21016  
ID AAU21016 standard; protein; 44 AA.  
XX  
AC AAU21016;  
XX  
DT 17-DEC-2001 (first entry)  
XX  
DE Human novel foetal antigen, SEQ ID NO 1260.  
XX  
KW Human; foetal tissue antigen; antiinflammatory; neuroprotective;  
KW immunomodulator; cardiovascular; cytostatic; nephrothropic;  
KW cardiovascular; autoimmune disease; rheumatoid arthritis;  
KW hyperproliferative disorder; breast neoplasm; cancer;  
KW cardiovascular disorder; cardiac arrest; cerebrovascular disorder;  
KW cerebral ischaemia; angiogenesis; nervous system disorder;  
KW Alzheimer's disease; infection; ocular disorder; corneal infection;  
KW wound healing; epithelial cell proliferation; food additive.  
XX  
OS Homo sapiens.  
XX  
FN WO200155312-A2.  
XX  
PD 02-AUG-2001.  
XX  
PF 17-JAN-2001; 2001WO-US001321.  
XX  
PR 31-JAN-2000; 2000US-0179065P.  
PR 04-FEB-2000; 2000US-0180628P.  
PR 24-FEB-2000; 2000US-0184564P.  
PR 02-MAR-2000; 2000US-0186350P.  
PR 16-MAR-2000; 2000US-0189874P.  
PR 17-MAR-2000; 2000US-0190076P.  
PR 18-APR-2000; 2000US-0198123P.  
PR 19-MAY-2000; 2000US-0205515P.  
PR 07-JUN-2000; 2000US-0209467P.  
PR 28-JUN-2000; 2000US-0214886P.  
PR 30-JUN-2000; 2000US-0215135P.  
PR 07-JUL-2000; 2000US-0216647P.  
PR 07-JUL-2000; 2000US-0216880P.  
PR 11-JUL-2000; 2000US-0217487P.  
PR 11-JUL-2000; 2000US-0217496P.  
PR 14-JUL-2000; 2000US-0218290P.  
PR 26-JUL-2000; 2000US-0220963P.  
PR 26-JUL-2000; 2000US-0220964P.  
PR 14-AUG-2000; 2000US-0224518P.  
PR 14-AUG-2000; 2000US-0224519P.  
PR 14-AUG-2000; 2000US-0225213P.  
PR 14-AUG-2000; 2000US-0225214P.  
PR 14-AUG-2000; 2000US-0225266P.  
PR 14-AUG-2000; 2000US-0225267P.  
PR 14-AUG-2000; 2000US-0225268P.  
PR 14-AUG-2000; 2000US-0225270P.  
PR 14-AUG-2000; 2000US-0225477P.  
PR 14-AUG-2000; 2000US-0225757P.  
PR 14-AUG-2000; 2000US-0225758P.  
PR 14-AUG-2000; 2000US-0225759P.  
PR 18-AUG-2000; 2000US-0226279P.  
PR 22-AUG-2000; 2000US-0226681P.  
PR 22-AUG-2000; 2000US-0226686P.  
PR 22-AUG-2000; 2000US-0227182P.  
PR 23-AUG-2000; 2000US-0227009P.  
PR 30-AUG-2000; 2000US-0228924P.  
PR 01-SEP-2000; 2000US-0229287P.  
PR 01-SEP-2000; 2000US-0229343P.  
PR 01-SEP-2000; 2000US-0229344P.  
PR 01-SEP-2000; 2000US-0229345P.  
PR 05-SEP-2000; 2000US-0229509P.  
PR 05-SEP-2000; 2000US-0229513P.  
PR 06-SEP-2000; 2000US-0230437P.  
PR 06-SEP-2000; 2000US-0230438P.  
PR 08-SEP-2000; 2000US-0231242P.  
PR 08-SEP-2000; 2000US-0231243P.  
PR 08-SEP-2000; 2000US-0231244P.  
PR 08-SEP-2000; 2000US-0231413P.  
PR 08-SEP-2000; 2000US-0231414P.  
PR 08-SEP-2000; 2000US-0232080P.  
PR 08-SEP-2000; 2000US-0232081P.  
PR 12-SEP-2000; 2000US-0231968P.  
PR 14-SEP-2000; 2000US-0232397P.  
PR 14-SEP-2000; 2000US-0232398P.  
PR 14-SEP-2000; 2000US-0232399P.  
PR 14-SEP-2000; 2000US-0232400P.  
PR 14-SEP-2000; 2000US-0232401P.  
PR 14-SEP-2000; 2000US-0233063P.  
PR 14-SEP-2000; 2000US-0233064P.  
PR 14-SEP-2000; 2000US-0233065P.  
PR 21-SEP-2000; 2000US-0234223P.  
PR 21-SEP-2000; 2000US-0234274P.  
PR 25-SEP-2000; 2000US-0234997P.  
PR 25-SEP-2000; 2000US-0234998P.  
PR 26-SEP-2000; 2000US-0235484P.  
PR 27-SEP-2000; 2000US-0235834P.  
PR 27-SEP-2000; 2000US-0235836P.  
PR 29-SEP-2000; 2000US-0236327P.  
PR 29-SEP-2000; 2000US-0236367P.  
PR 29-SEP-2000; 2000US-0236368P.  
PR 29-SEP-2000; 2000US-0236369P.  
PR 29-SEP-2000; 2000US-0236370P.  
PR 02-OCT-2000; 2000US-0236802P.  
PR 02-OCT-2000; 2000US-0237037P.  
PR 02-OCT-2000; 2000US-0237038P.  
PR 02-OCT-2000; 2000US-0237039P.  
PR 13-OCT-2000; 2000US-0237040P.  
PR 13-OCT-2000; 2000US-0239935P.  
PR 13-OCT-2000; 2000US-0239937P.  
PR 20-OCT-2000; 2000US-0240960P.  
PR 20-OCT-2000; 2000US-0241221P.  
PR 20-OCT-2000; 2000US-0241785P.  
PR 20-OCT-2000; 2000US-0241786P.

PR 20-OCT-2000; 2000US-0241787P.  
 PR 20-OCT-2000; 2000US-0241808P.  
 PR 20-OCT-2000; 2000US-0241809P.  
 PR 20-OCT-2000; 2000US-0241826P.  
 PR 01-NOV-2000; 2000US-0244617P.  
 PR 08-NOV-2000; 2000US-0246474P.  
 PR 08-NOV-2000; 2000US-0246475P.  
 PR 08-NOV-2000; 2000US-0246476P.  
 PR 08-NOV-2000; 2000US-0246477P.  
 PR 08-NOV-2000; 2000US-0246478P.  
 PR 08-NOV-2000; 2000US-0246523P.  
 PR 08-NOV-2000; 2000US-0246524P.  
 PR 08-NOV-2000; 2000US-0246525P.  
 PR 08-NOV-2000; 2000US-0246526P.  
 PR 08-NOV-2000; 2000US-0246527P.  
 PR 08-NOV-2000; 2000US-0246528P.  
 PR 08-NOV-2000; 2000US-0246532P.  
 PR 08-NOV-2000; 2000US-0246609P.  
 PR 08-NOV-2000; 2000US-0246610P.  
 PR 08-NOV-2000; 2000US-0246611P.  
 PR 08-NOV-2000; 2000US-0246613P.  
 PR 17-NOV-2000; 2000US-0249207P.  
 PR 17-NOV-2000; 2000US-0249208P.  
 PR 17-NOV-2000; 2000US-0249209P.  
 PR 17-NOV-2000; 2000US-0249210P.  
 PR 17-NOV-2000; 2000US-0249211P.  
 PR 17-NOV-2000; 2000US-0249212P.  
 PR 17-NOV-2000; 2000US-0249213P.  
 PR 17-NOV-2000; 2000US-0249214P.  
 PR 17-NOV-2000; 2000US-0249215P.  
 PR 17-NOV-2000; 2000US-0249216P.  
 PR 17-NOV-2000; 2000US-0249217P.  
 PR 17-NOV-2000; 2000US-0249218P.  
 PR 17-NOV-2000; 2000US-0249244P.  
 PR 17-NOV-2000; 2000US-0249245P.  
 PR 17-NOV-2000; 2000US-0249264P.  
 PR 17-NOV-2000; 2000US-0249265P.  
 PR 17-NOV-2000; 2000US-0249297P.  
 PR 17-NOV-2000; 2000US-0249299P.  
 PR 17-NOV-2000; 2000US-0249300P.  
 PR 01-DEC-2000; 2000US-0250160P.  
 PR 01-DEC-2000; 2000US-0250391P.  
 PR 05-DEC-2000; 2000US-0251030P.  
 PR 05-DEC-2000; 2000US-0251388P.  
 PR 05-DEC-2000; 2000US-0256719P.  
 PR 05-DEC-2000; 2000US-0251479P.  
 PR 08-DEC-2000; 2000US-0251856P.  
 PR 08-DEC-2000; 2000US-0251868P.  
 PR 08-DEC-2000; 2000US-0251869P.  
 PR 08-DEC-2000; 2000US-0251989P.  
 PR 08-DEC-2000; 2000US-0251990P.  
 PR 11-DEC-2000; 2000US-0254097P.  
 PR 05-JAN-2001; 2001US-0259678P.  
 PR (HUMA-) HUMAN GENOME SCI INC.  
 XX  
 PI Rosen CA, Barash SC, Ruben SM;  
 XX  
 DR WPI; 2001-488792/53.  
 DR N-PSDB; AAS33836.  
 XX  
 PT New polynucleotides and polypeptides for diagnosing, treating, preventing  
 PT or prognosing e.g. diseases or disorders of the nervous, musculoskeletal,  
 PT excretory, gastrointestinal, reproductive, and respiratory systems.  
 XX  
 PS Claim 11; SEQ ID NO 1260; 642pp; English.  
 XX  
 CC The invention relates to novel nucleic acids encoding novel human foetal  
 CC antigens. The nucleic acids and proteins are used to prevent, treat (e.g.  
 CC by gene therapy) or ameliorate a medical condition in e.g. humans, mice,  
 CC rabbits, goats, horses, cats, dogs, chickens or sheep. They are also used  
 CC in diagnosing a pathological condition or susceptibility to a  
 CC pathological condition. The antibodies to the antigens can also be used

CC in alleviating symptoms associated with the disorders and in diagnostic  
 CC immunoassays e.g. radioimmunoassays or enzyme linked immunosorbent assays  
 CC (ELISA). Disorders which are diagnosed or treated include autoimmune  
 CC diseases e.g. rheumatoid arthritis, hyperproliferative disorders e.g.  
 CC neoplasms of the breast or liver, cardiovascular disorders e.g. cardiac  
 CC arrest, cerebrovascular disorders e.g. cerebral ischaemia, angiogenesis,  
 CC nervous system disorders e.g. Alzheimer's disease, infections caused by  
 CC bacteria, viruses and fungi and ocular disorders e.g. corneal infection.  
 CC The polypeptides can also be used to aid wound healing and epithelial  
 CC cell proliferation, to prevent skin aging due to sunburn, to maintain  
 CC organs before transplantation, for supporting cell culture of primary  
 CC tissues, to regenerate tissues and in chemotaxis. The polypeptides can  
 CC also be used as a food additive or preservative to increase or decrease  
 CC storage capabilities, fat content, lipid, protein, carbohydrate,  
 CC vitamins, minerals, cofactors and other nutritional components. Numerous  
 CC examples of diseases and disorders treated by the nucleic acids and  
 CC proteins are given in the specification. The present sequence represents  
 CC a foetal antigen of the invention. Note: The sequence data for this  
 CC patent did not form part of the printed specification, but was obtained

Query Match 32.0%; Score 31; DB 4; Length 44;  
 Best Local Similarity 75.0%; Pred. No. 8.5e+02;  
 Matches 6; Conservative 1; Mismatches 1; Indels 0; Gaps 0;  
 QY 5 LNSKIAFK 12  
 Db 24 LNSKLJFK 31  
 ||||: ||  
 ||||: ||

RESULT 58  
 ABG46440  
 ID ABG46440 standard; peptide; 44 AA.  
 XX  
 AC ABG46440;  
 XX  
 DT 19-AUG-2002 (first entry)  
 XX  
 DE Human peptide encoded by genome-derived single exon probe SEQ ID 36105.  
 XX  
 KW Human; single exon probe; asthma; lung cancer; COPD; ILD;  
 KW chronic obstructive pulmonary disease; interstitial lung disease;  
 KW familial idiopathic pulmonary fibrosis; neurofibromatosis;  
 KW tuberous sclerosis; Gaucher's disease; Niemann-Pick disease;  
 KW Hermansky-Pudlak syndrome; sarcoidosis; pulmonary haemosiderosis;  
 KW pulmonary histiocytosis; lymphangioleiomyomatosis; Karagener syndrome;  
 KW pulmonary alveolar proteinosis; fibrocystic pulmonary dysplasia;  
 KW primary ciliary dyskinesia; pulmonary hypertension;  
 KW hyaline membrane disease.  
 XX  
 OS Homo sapiens.  
 XX  
 PN WO200186003-A2.  
 XX  
 PD 15-NOV-2001.  
 XX  
 PF 30-JAN-2001; 2001WO-US000665.  
 XX  
 PR 04-FEB-2000; 2000US-0180312P.  
 PR 26-MAY-2000; 2000US-0207456P.  
 PR 30-JUN-2000; 2000US-00608408.  
 PR 03-AUG-2000; 2000US-00632366.  
 PR 21-SEP-2000; 2000US-0234687P.  
 PR 07-SEP-2000; 2000US-0236359P.  
 PR 04-OCT-2000; 2000GB-00024263.  
 XX  
 PA (MOLE-) MOLECULAR DYNAMICS INC.  
 XX  
 PI Penn SG, Hanzel DK, Chen W, Rank DR;  
 XX  
 DR WPI; 2002-114183/15.  
 XX  
 PT Spatially-addressable set of single exon nucleic acid probes, used to  
 PT measure gene expression in human lung samples.

```
XX PS Claim 27; SEQ ID NO 36105; 634pp; English.
XX CC
XX CC The invention relates to a spatially-addressable set of single exon
XX CC nucleic acid probes for measuring gene expression in a sample derived
XX CC from human lung comprising single exon nucleic acid probes having one of
XX CC 12614 nucleic acid sequences mentioned in the specification, or their
XX CC complements or the 12387 open reading frames derived from the 12614
XX CC probes. Also included are a microarray comprising the novel set of probes
XX CC ; the novel set of probes which hybridise at high stringency to a nucleic
XX CC acid expressed in the human lung; measuring gene expression in a sample
XX CC derived from human lung, comprising (a) contacting the array with a
XX CC collection of detectably labeled nucleic acids derived from human lung
XX CC mRNA, and (b) measuring the label detectably bound to each probe of the
XX CC array; identifying exons in a eukaryotic genome, comprising (a)
XX CC algorithmically predicting at least one exon from genomic sequences of
XX CC the eukaryote; and (b) detecting specific hybridisation of detectably
XX CC labeled nucleic acids from eukaryote lung mRNA, to a single exon probe,
XX CC having a fragment identical to the predicted exon, the probe is included
XX CC in the above mentioned microarray; assigning exons to a single gene,
XX CC comprising (a) identifying exons from genomic sequence by the method
XX CC above and (b) measuring the expression of each of the exons in several
XX CC tissues and/or cell types using hybridisation to a single exon
XX CC microarrays having a probe with the exon, where a common pattern of
XX CC expression of the exons in the tissues and/or cell types indicates that
XX CC the exons should be assigned to a single gene; a peptide comprising one
XX CC of 12011 sequences, mentioned in the specification, or encoded by the
XX CC probes/open reading frames (ORF). The probes are used for gene expression
XX CC analysis, and for identifying exons in a gene, particularly using human
XX CC lung derived mRNA and for the study of lung diseases such as asthma, lung
XX CC cancer, chronic obstructive pulmonary disease (COPD), interstitial lung
XX CC disease (ILD), familial idiopathic pulmonary fibrosis, neurofibromatosis,
XX CC tuberous sclerosis, Gaucher's disease, Niemann-Pick disease, Hermansky-
XX CC Pudlak syndrome, sarcoidosis, pulmonary haemosiderosis, pulmonary
XX CC histiocytosis, lymphangioleiomyomatosis, pulmonary alveolar proteinosis,
XX CC Karagenex syndrome, fibrocystic pulmonary dysplasia, primary ciliary
XX CC dyskinesia, pulmonary hypertension and hyaline membrane disease. The
XX CC present sequence is a peptide/protein encoded by a single exon probe of
XX CC the invention. Note: The sequence data for this patent did not form part
XX CC of the printed specification, but was obtained in electronic format
XX CC directly from WIPO at ftp.wipo.int/pub/published_pct_sequences
XX SQ Sequence 44 AA;

Query Match 32.0%; Score 31; DB 5; Length 44;
Best Local Similarity 33.3%; Pred. No. 8.5e+02;
Matches 8; Conservative 3; Mismatches 5; Indels 8; Gaps 1;

QY 3 NHLNSKIA-----FKIVGQEP 18
DB :|:|:|:|:|:|:|:|:|
9 DHINLVKAGQGSVVQFKIKRHTP 32

RESULT 59
ADF70009
ID ADF70009 standard; protein; 44 AA.
XX AC ADF70009;
XX DT 12-FEB-2004 (first entry)
XX DE AcmA-type homologue amino acid sequence.
XX KW delivery; targeting system; AcmA-type anchor protein; solid tumour;
XX KW health; medical; agricultural; cosmetic; controlled release.
XX OS Lactococcus lactis.
XX PN WO2003084508-A1.
XX PD 16-OCT-2003.
XX PF 04-APR-2003; 2003WO-NL000256.
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XX PR 04-APR-2002; 2002EP-00076316.
XX PR 04-APR-2002; 2002US-0369927P.
XX PR 05-APR-2002; 2002US-0370485P.
XX PR 20-DEC-2002; 2002EP-00080481.
XX PA (NANO-) APPLIED NANOSYSTEMS BV.
XX FI Friesen RHE, Leenhouts CJ, Hektor HJ, Van Esch JH, Heeres A;
XX Robilliard GT;
XX DR WPI; 2003-877005/81.
XX PT Vehicle for targeted delivery of therapeutic or diagnostic agents,
XX PT includes protein anchor and system for inducing availability of the
XX PT agent.
XX PS Example 3; Page 191; 303pp; English.
XX CC The present invention describes a vehicle (A) for delivering a substance
XX CC (I) to a predetermined site, which comprises (I); a system for inducing
XX CC availability of at least one compartment of (A) towards the exterior;
XX CC and, as targeting system for directing (A) to the site, an AcmA-type
XX CC anchor protein (II). (A) are used for delivery of diagnostic and
XX CC therapeutic agents to predetermined sites in the body, particularly
XX CC joints or solid tumours but can be used more generally for health,
XX CC medical, agricultural and cosmetic applications. (A) significantly
XX CC increases the half-life of peptides in the circulation and, by providing
XX CC controlled release, ensures relatively high bioavailability, allowing
XX CC therapeutic use of agents that would otherwise be too toxic for systemic
XX CC administration. The native AcmA peptide targets Gram-positive bacteria
XX CC but its homologues can be engineered to have different selectivity. The
XX CC present sequence is used in the exemplification of the present invention.
XX SQ Sequence 44 AA;

Query Match 32.0%; Score 31; DB 7; Length 44;
Best Local Similarity 66.7%; Pred. No. 8.5e+02;
Matches 6; Conservative 0; Mismatches 3; Indels 0; Gaps 0;

QY 3 NHLNSKIAP 11
DB :|:|:|:|:|:|
28 NHLNSDTIP 36

RESULT 60
AAB16298
ID AAB16298 standard; protein; 47 AA.
XX AC AAB16298;
XX DT 31-OCT-2000 (first entry)
XX DE Eucalyptus grandis AGP protein sequence SEQ ID NO:84.
XX KW Eucalyptus grandis; pinus radiata; Monterey pine; modification;
XX KW plant cell wall; polysaccharide; polysaccharide biosynthetic pathway;
XX KW transgenic plant.
XX OS Eucalyptus grandis.
XX PN WO200022092-A2.
XX PD 20-APR-2000.
XX PF 08-OCT-1999; 99WO-NZ000169.
XX PR 13-OCT-1998; 98US-00170862.
XX PR 11-AUG-1999; 99US-0148426P.
XX PA (GENE-) GENESIS RES & DEV CORP LTD.
XX PA (FLET-) FLETCHER CHALLENGE FORESTS LTD.
XX
```

PI Bloksberg LN;  
 XX WPI; 2000-339328/29.  
 DR N-PSDB; AAA67105.  
 XX  
 PT New genes encoding proteins involved in a plant polysaccharide  
 PT biosynthetic pathway, useful for modulating or altering the  
 PT polysaccharide content, composition or structure of the plant.  
 XX  
 PS Claim 17; Page 78; 301pp; English.  
 XX  
 CC The present invention describes isolated polynucleotides (PN) comprising  
 CC a sequence selected from one of 835 nucleotide sequences given in  
 CC AAA67073 to AAA67907, their (reverse) complements, sequences producing an  
 CC Expectation (E) value of 0.01 or less compared to the 835 sequences, the  
 CC sequences at least 50% identical to them, 200, 100, 40 or 20-mers of the  
 CC 835 sequences or sequences that are degenerately equivalent or allelic to  
 CC the 835 sequences. The polynucleotides are used to modify the activity of  
 CC a polypeptide involved in a polysaccharide biosynthetic pathway in the  
 CC plant. They are especially used to modulate or alter the polysaccharide  
 CC content, composition or structure of the plant. AAB16268 to AAB16340 are  
 CC proteins encoded by some of the polynucleotide sequence given in the  
 CC present invention  
 XX  
 SQ Sequence 47 AA;  
 Query Match 32.0%; Score 31; DB 3; Length 47;  
 Best Local Similarity 50.0%; Pred. No. 9.2e+02;  
 Matches 7; Conservative 3; Mismatches 4; Indels 0; Gaps 0;  
 Qy 5 LNSKIAFKIVSQEP 18  
 Db 4 LDSADAFKSVRRDP 17  
 RESULT 61  
 ADJ12233  
 ID ADJ12233 standard; protein; 47 AA.  
 AC ADJ12233;  
 XX  
 DT 20-MAY-2004 (first entry)  
 DE  
 XX Human secreted protein SeqID 87.  
 KW human; secreted; cancer; haematopoietic disease; anaemia;  
 KW multiple myeloma; reproductive system disorder; prostatitis;  
 KW inguinal hernia; musculoskeletal disease; systemic lupus erythematosus;  
 KW gout; cardiovascular disease; arrhythmia; hypernatraemia; fetal disease;  
 KW fetal alcohol syndrome; Down's syndrome; excretory disease;  
 KW urinary incontinence; renal disorder; neural; sensory disease;  
 KW Alzheimer's disease; meningitis; respiratory disease; emphysema;  
 KW occupational lung disease; endocrine disease; diabetes;  
 KW glomerulonephritis; digestive disease; portal hypertension;  
 KW irritable bowel syndrome; epithelial disease; scleroderma;  
 KW epidermolysis bullosa; cytostatic; antianemic; antiarthritic;  
 KW antiasthmatic; anti-HIV; immunosuppressive; antiinflammatory;  
 KW antipsoriatic; antibacterial; osteopathic; dermatological; antigout;  
 KW immunomodulator; antiarrhythmic; cardiant; nootropic; antileptic;  
 KW nephrotropic; uropathic; neuroprotective; antiparkinsonian; tranquilizer;  
 KW antidiabetic; anabolic; hypertensive; vulnary.  
 XX  
 OS Homo sapiens.  
 XX  
 XX US2004010132-A1.  
 PD 15-JAN-2004.  
 XX  
 PF 30-OCT-2001; 2001US-00984429.  
 XX  
 PR 09-OCT-1997; 97US-0061463P.  
 PR 09-OCT-1997; 97US-0061527P.  
 PR 09-OCT-1997; 97US-0061529P.

PR 09-OCT-1997; 97US-0061532P.  
 PR 09-OCT-1997; 97US-0061536P.  
 PR 09-OCT-1997; 97US-0071498P.  
 PR 08-OCT-1998; 98WO-US021142.  
 PR 08-APR-1999; 99US-00288143.  
 PR 01-NOV-2000; 2000US-0244591P.  
 XX  
 PA (ROSE/) ROSEN C A.  
 PA (BREW/) BREWER L A.  
 PA (DUAN/) DUAN R D.  
 PA (RUBE/) RUBEN S M.  
 PA (FLOR/) FLORENCE K A.  
 PA (GREE/) GREENE J M.  
 PA (YOUN/) YOUNG P E.  
 PA (FERR/) FERRIE A M.  
 PA (YUGG/) YU G.  
 PA (FLOR/) FLORENCE C.  
 PA (EBNE/) EBNER R.  
 PA (OLSE/) OLSEN H.  
 XX  
 FI Rosen CA, Brewer LA, Duan RD, Ruben SM, Florence KA, Greene JM;  
 PI Young PE, Ferrie AM, Yu G, Florence C, Ebner R, Olsen H;  
 XX  
 DR WPI; 2004-090518/09.  
 DR N-PSDB; ADJ12177.  
 XX  
 PT New isolated nucleic acids and polypeptides, useful for diagnosing,  
 PT treating, preventing or ameliorating diseases or disorders e.g. cancer,  
 PT anaemia, arthritis, asthma, inflammatory bowel disease or Alzheimer's  
 PT disease.  
 XX  
 PS Claim 11; SEQ ID NO 87; 286pp; English.  
 XX  
 CC This invention relates to novel polynucleotides encoding human secreted  
 CC proteins. Specifically, it refers to the vectors, host cells, recombinant  
 CC and synthetic methods for producing human polynucleotides, polypeptides  
 CC and antibodies. Furthermore, it relates to screening methods to identify  
 CC agonists and antagonists that can be used to inhibit or enhance the  
 CC production and function of the secreted proteins. The present invention  
 CC describes these compositions as useful for diagnosing, treating or  
 CC preventing disorders such as cancer, haematopoietic diseases including  
 CC anaemia and multiple myeloma, reproductive system disorders including  
 CC prostatitis and inguinal hernia, musculoskeletal diseases including  
 CC systemic lupus erythematosus and gout, cardiovascular disease including  
 CC arrhythmia and hypernatraemia, mixed fetal diseases including fetal  
 CC alcohol syndrome and Down's syndrome, excretory diseases including  
 CC urinary incontinence and renal disorders, neural or sensory disease  
 CC including Alzheimer's disease and meningitis, respiratory disease  
 CC including emphysema and occupational lung disease, endocrine diseases  
 CC including diabetes and glomerulonephritis, digestive diseases including  
 CC portal hypertension and irritable bowel syndrome and connective tissue or  
 CC epithelial diseases including scleroderma and epidermolysis bullosa. As  
 CC such, there are various activities such as cytostatic, antianemic,  
 CC antiarthritic, antiasthmatic, anti-HIV, immunosuppressive,  
 CC antiinflammatory, antipsoriatic, antibacterial, osteopathic,  
 CC dermatological, antigout, immunomodulator, antiarrhythmic, cardiant,  
 CC nootropic, antileptic, neuroprotective, uropathic, neuroprotective,  
 CC antiparkinsonian, tranquilizer, antidiabetic, anabolic, hypertensive and  
 CC vulnary. This polypeptide is a human secreted protein of the invention.  
 CC NOTE: This sequence does not appear in the printed specification but has  
 CC been obtained in electronic format from the US patent office at the  
 CC following web site [www.seqdata.uspto.gov/sequence.html](http://www.seqdata.uspto.gov/sequence.html); Document ID:  
 CC 20040010132.  
 XX  
 SQ Sequence 47 AA;  
 Query Match 32.0%; Score 31; DB 8; Length 47;  
 Best Local Similarity 45.5%; Pred. No. 9.2e+02;  
 Matches 5; Conservative 4; Mismatches 2; Indels 0; Gaps 0;  
 Qy 9 IAFKIVSQEPA 19  
 Db 19 VAFRLTNQIPA 29

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RESULT 62
AAAY14431
ID  AAAY14431 standard; protein; 48 AA.
XX
XX
AC  AAAY14431;
XX
XX
DT  17-AUG-1999 (first entry)
XX
XX
DE  Human secreted protein encoded by gene 21 clone HRDED19.
XX
XX
KW  Human; secreted protein; fusion protein; gene therapy; protein therapy;
KW  diagnosis; tissue; cancer; tumour; neurodegenerative disorder; leukaemia;
KW  developmental abnormality; foetal deficiency; blood; allergy; renal;
KW  immune system; asthma; lymphocytic disease; brain; hepatic; lymphoma;
KW  inflammation; ischaemic shock; Alzheimer's disease; restenosis; AIDS;
KW  cognitive disorder; schizophrenia; prostate; obesity; osteoclast; thymus;
KW  osteoporosis; arthritis; testis; lung; thyroiditis; thyroid; digestion;
KW  endocrine; metabolism; regulation; malabsorption; gastritis; neoplasm.
XX
XX
OS  Homo sapiens.
XX
XX
FN  WO9919339-A1.
XX
XX
PD  22-APR-1999.
XX
XX
PF  08-OCT-1998; 98WO-US021142.
XX
XX
PR  09-OCT-1997; 97US-0061463P.
XX
PR  09-OCT-1997; 97US-0061527P.
XX
PR  09-OCT-1997; 97US-0061529P.
XX
PR  09-OCT-1997; 97US-0061532P.
XX
PR  09-OCT-1997; 97US-0061536P.
XX
PR  09-OCT-1997; 97US-0071498P.
XX
XX
PA  (HUMA-) HUMAN GENOME SCI INC.
XX
XX
PI  Brewer LA, Olsen HS, Duan R, Ebner R, Rosen CA, Ruben SM;
PI  Florence KA, Young FE, Greene JW, Yu G, Ferrie AM, Florence C;
XX
XX
WPI; 1999-277587/23.
DR  N-PSDB; AAX79031.
XX
XX
PT  New isolated human genes and the secreted polypeptides they encode.
XX
XX
PS  Claim 11; Page 192; 226pp; English.
XX
XX
This sequence represents a secreted human protein encoded by the gene
CC  clone detailed in the descriptor line. The gene can be used to generate
CC  fusion proteins by linking to the gene to a human immunoglobulin Fc
CC  portion (e.g. AAX79002) for increasing the stability of the fused protein
CC  as compared to the human protein only. The invention relates to 53 novel
CC  genes and their fragments (nucleic acid sequences: AAX79011-X79064; amino
CC  acid sequences AAAY1441-Y14464) which are useful for preventing, treating
CC  or ameliorating medical conditions e.g. by protein or gene therapy. Also,
CC  pathological conditions can be diagnosed by determining the amount of the
CC  new polypeptides in a sample or by determining the presence of mutations
CC  in the new polynucleotides. Specific uses are described for each of the
CC  53 polynucleotides, based on which tissues they are most highly expressed
CC  in (see AAX79011 for described uses)
XX
XX
SQ  Sequence 48 AA;
XX
Query Match 32.0%; Score 31; DB 2; Length 48;
Best Local Similarity 45.5%; Pred. No. 9.4e+02;
Matches 5; Conservative 4; Mismatches 2; Indels 0; Gaps 0;
XX
QY 9 IAFKIVSQEPA 19
:|:|:|:|
DB 19 VAFRLTNQIPA 29
:|:|:|:|
XX
RESULT 63
ADE86990
ID  ADE86990 standard; protein; 49 AA.
XX
XX
AC  ADE86990;
XX
XX
DT  29-JAN-2004 (first entry)
XX
XX
DE  Human pancreatic cell protein sequence SeqID450.
XX
XX
KW  neoplastic pancreatic cell; pancreatic cell; pancreatic cancer;
KW  cancer death; cytostatic; vaccine; gene therapy;
KW  non-cancerous pancreas disease; human.
XX
XX
OS  Homo sapiens.
XX
XX
FN  WO2003060145-A2.
XX
XX
PD  24-JUL-2003.
XX
XX
PF  19-DEC-2002; 2002WO-US040655.
XX
XX
PR  21-DEC-2001; 2001US-0342768P.
XX
XX
PA  (DIAD-) DIADEXUS INC.
XX
XX
PI  Sun Y, Liu C;
XX
XX
WPI; 2003-587286/55.
DR  N-PSDB; ADE87263.
XX
XX
PT  New pancreatic specific nucleic acid molecule or protein for diagnosing,
PT  staging, imaging, monitoring, preventing or treating pancreatic cancer or
PT  non-cancerous disease states of the pancreas.
XX
XX
PS  Claim 12; SEQ ID NO 450; 635pp; English.
XX
XX
This invention relates to novel nucleic acids and proteins present in
CC  normal and neoplastic pancreatic cells. Pancreatic cancer is a common
CC  cause of cancer death worldwide, therefore accurate methods of diagnosis
CC  and treatment are required. Compounds which modulate the proteins of the
CC  invention may have cytostatic activity and the protein and DNA sequences
CC  of the invention may be useful for the development of a vaccine or in
CC  gene therapy. The composition and methods are useful in diagnosing,
CC  staging, imaging, monitoring, preventing or treating pancreatic cancer
CC  and non-cancerous disease states of the pancreas. The present sequence is
CC  that of a human pancreatic protein of the invention.
XX
XX
SQ  Sequence 49 AA;
XX
Query Match 32.0%; Score 31; DB 7; Length 49;
Best Local Similarity 50.0%; Pred. No. 9.7e+02;
Matches 5; Conservative 3; Mismatches 2; Indels 0; Gaps 0;
XX
QY 2 PNHLNSKTIAP 11
:|:|:|:|
DB 2 PNYLHSSLPP 11
:|:|:|:|
XX
RESULT 64
AAAU17741
ID  AAU17741 standard; protein; 50 AA.
XX
XX
AC  AAU17741;
XX
XX
DT  07-NOV-2001 (first entry)
XX
XX
DE  Novel human respiratory antigen #57.
XX
XX
KW  Human; respiratory antigen; respiratory disorder; throat disorder;
KW  lung disorder; nose disorder; lung cancer; gene therapy; cytostatic;
KW  anti allergic; anti asthmatic; anti inflammatory; olfactory;
KW  respiratory active.

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XX  
OS Homo sapiens.  
XX  
PN WO200155448-A1.  
XX  
PD 02-AUG-2001.  
XX  
PF 17-JAN-2001; 2001WO-US001333.  
XX  
XX 31-JAN-2000; 2000US-0179065P.  
PR 04-FEB-2000; 2000US-0180628P.  
PR 24-FEB-2000; 2000US-0184664P.  
PR 02-MAR-2000; 2000US-0186350P.  
PR 16-MAR-2000; 2000US-0189874P.  
PR 17-MAR-2000; 2000US-0190076P.  
PR 18-APR-2000; 2000US-0198123P.  
PR 19-MAY-2000; 2000US-0205515P.  
PR 07-JUN-2000; 2000US-0209467P.  
PR 28-JUN-2000; 2000US-0214886P.  
PR 30-JUN-2000; 2000US-0215135P.  
PR 07-JUL-2000; 2000US-0216647P.  
PR 07-JUL-2000; 2000US-0216880P.  
PR 11-JUL-2000; 2000US-0217487P.  
PR 11-JUL-2000; 2000US-0217496P.  
PR 14-JUL-2000; 2000US-0218230P.  
PR 26-JUL-2000; 2000US-0220963P.  
PR 26-JUL-2000; 2000US-0220964P.  
PR 14-AUG-2000; 2000US-0224518P.  
PR 14-AUG-2000; 2000US-0224519P.  
PR 14-AUG-2000; 2000US-0225213P.  
PR 14-AUG-2000; 2000US-0225214P.  
PR 14-AUG-2000; 2000US-0225266P.  
PR 14-AUG-2000; 2000US-0225267P.  
PR 14-AUG-2000; 2000US-0225268P.  
PR 14-AUG-2000; 2000US-0225270P.  
PR 14-AUG-2000; 2000US-0225447P.  
PR 14-AUG-2000; 2000US-0225757P.  
PR 14-AUG-2000; 2000US-0225758P.  
PR 14-AUG-2000; 2000US-0225759P.  
PR 18-AUG-2000; 2000US-0226279P.  
PR 22-AUG-2000; 2000US-0226661P.  
PR 22-AUG-2000; 2000US-0226686P.  
PR 22-AUG-2000; 2000US-0227182P.  
PR 23-AUG-2000; 2000US-0227009P.  
PR 30-AUG-2000; 2000US-0228924P.  
PR 01-SEP-2000; 2000US-0229387P.  
PR 01-SEP-2000; 2000US-0229343P.  
PR 01-SEP-2000; 2000US-0229344P.  
PR 01-SEP-2000; 2000US-0229345P.  
PR 05-SEP-2000; 2000US-0229509P.  
PR 05-SEP-2000; 2000US-0229513P.  
PR 06-SEP-2000; 2000US-0230437P.  
PR 06-SEP-2000; 2000US-0230438P.  
PR 08-SEP-2000; 2000US-0231242P.  
PR 08-SEP-2000; 2000US-0231243P.  
PR 08-SEP-2000; 2000US-0231244P.  
PR 08-SEP-2000; 2000US-0231413P.  
PR 08-SEP-2000; 2000US-0231414P.  
PR 08-SEP-2000; 2000US-0232080P.  
PR 08-SEP-2000; 2000US-0232081P.  
PR 12-SEP-2000; 2000US-0231968P.  
PR 14-SEP-2000; 2000US-0232397P.  
PR 14-SEP-2000; 2000US-0232398P.  
PR 14-SEP-2000; 2000US-0232399P.  
PR 14-SEP-2000; 2000US-0232400P.  
PR 14-SEP-2000; 2000US-0232401P.  
PR 14-SEP-2000; 2000US-0233063P.  
PR 14-SEP-2000; 2000US-0233064P.  
PR 14-SEP-2000; 2000US-0233065P.  
PR 21-SEP-2000; 2000US-0234223P.  
PR 21-SEP-2000; 2000US-0234274P.  
PR 23-SEP-2000; 2000US-0234957P.  
PR 25-SEP-2000; 2000US-0234958P.  
XX  
XX 26-SEP-2000; 2000US-0235484P.  
PR 27-SEP-2000; 2000US-0235834P.  
PR 27-SEP-2000; 2000US-0235836P.  
PR 29-SEP-2000; 2000US-0236327P.  
PR 29-SEP-2000; 2000US-0236367P.  
PR 29-SEP-2000; 2000US-0236368P.  
PR 29-SEP-2000; 2000US-0236369P.  
PR 29-SEP-2000; 2000US-0236370P.  
PR 02-OCT-2000; 2000US-0236802P.  
PR 02-OCT-2000; 2000US-0237037P.  
PR 02-OCT-2000; 2000US-0237038P.  
PR 02-OCT-2000; 2000US-0237039P.  
PR 13-OCT-2000; 2000US-0237040P.  
PR 13-OCT-2000; 2000US-0239355P.  
PR 13-OCT-2000; 2000US-0239357P.  
PR 20-OCT-2000; 2000US-0240960P.  
PR 20-OCT-2000; 2000US-0241221P.  
PR 20-OCT-2000; 2000US-0241785P.  
PR 20-OCT-2000; 2000US-0241786P.  
PR 20-OCT-2000; 2000US-0241787P.  
PR 20-OCT-2000; 2000US-0241808P.  
PR 20-OCT-2000; 2000US-0241809P.  
PR 01-NOV-2000; 2000US-0244617P.  
PR 08-NOV-2000; 2000US-0246474P.  
PR 08-NOV-2000; 2000US-0246475P.  
PR 08-NOV-2000; 2000US-0246476P.  
PR 08-NOV-2000; 2000US-0246477P.  
PR 08-NOV-2000; 2000US-0246478P.  
PR 08-NOV-2000; 2000US-0246523P.  
PR 08-NOV-2000; 2000US-0246524P.  
PR 08-NOV-2000; 2000US-0246525P.  
PR 08-NOV-2000; 2000US-0246526P.  
PR 08-NOV-2000; 2000US-0246527P.  
PR 08-NOV-2000; 2000US-0246528P.  
PR 08-NOV-2000; 2000US-0246532P.  
PR 08-NOV-2000; 2000US-0246609P.  
PR 08-NOV-2000; 2000US-0246610P.  
PR 08-NOV-2000; 2000US-0246611P.  
PR 08-NOV-2000; 2000US-0246613P.  
PR 17-NOV-2000; 2000US-0249207P.  
PR 17-NOV-2000; 2000US-0249208P.  
PR 17-NOV-2000; 2000US-0249209P.  
PR 17-NOV-2000; 2000US-0249210P.  
PR 17-NOV-2000; 2000US-0249211P.  
PR 17-NOV-2000; 2000US-0249212P.  
PR 17-NOV-2000; 2000US-0249213P.  
PR 17-NOV-2000; 2000US-0249214P.  
PR 17-NOV-2000; 2000US-0249215P.  
PR 17-NOV-2000; 2000US-0249216P.  
PR 17-NOV-2000; 2000US-0249217P.  
PR 17-NOV-2000; 2000US-0249218P.  
PR 17-NOV-2000; 2000US-0249244P.  
PR 17-NOV-2000; 2000US-0249245P.  
PR 17-NOV-2000; 2000US-0249264P.  
PR 17-NOV-2000; 2000US-0249265P.  
PR 17-NOV-2000; 2000US-0249297P.  
PR 17-NOV-2000; 2000US-0249299P.  
PR 17-NOV-2000; 2000US-0249300P.  
PR 01-DEC-2000; 2000US-0250160P.  
PR 01-DEC-2000; 2000US-0250391P.  
PR 05-DEC-2000; 2000US-0251030P.  
PR 05-DEC-2000; 2000US-0251988P.  
PR 06-DEC-2000; 2000US-0256719P.  
PR 08-DEC-2000; 2000US-0251479P.  
PR 08-DEC-2000; 2000US-0251856P.  
PR 08-DEC-2000; 2000US-0251868P.  
PR 08-DEC-2000; 2000US-0251869P.  
PR 08-DEC-2000; 2000US-0251989P.  
PR 11-DEC-2000; 2000US-0251990P.  
PR 05-JAN-2001; 2001US-0259678P.  
XX  
XX



PR 20-OCT-2000; 2000US-0240960P.  
 PR 20-OCT-2000; 2000US-0241221P.  
 PR 20-OCT-2000; 2000US-0241785P.  
 PR 20-OCT-2000; 2000US-0241786P.  
 PR 20-OCT-2000; 2000US-0241787P.  
 PR 20-OCT-2000; 2000US-0241808P.  
 PR 20-OCT-2000; 2000US-0241809P.  
 PR 20-OCT-2000; 2000US-0241826P.  
 PR 01-NOV-2000; 2000US-0244617P.  
 PR 08-NOV-2000; 2000US-0246474P.  
 PR 08-NOV-2000; 2000US-0246475P.  
 PR 08-NOV-2000; 2000US-0246476P.  
 PR 08-NOV-2000; 2000US-0246477P.  
 PR 08-NOV-2000; 2000US-0246478P.  
 PR 08-NOV-2000; 2000US-0246523P.  
 PR 08-NOV-2000; 2000US-0246524P.  
 PR 08-NOV-2000; 2000US-0246525P.  
 PR 08-NOV-2000; 2000US-0246526P.  
 PR 08-NOV-2000; 2000US-0246527P.  
 PR 08-NOV-2000; 2000US-0246528P.  
 PR 08-NOV-2000; 2000US-0246532P.  
 PR 08-NOV-2000; 2000US-0246609P.  
 PR 08-NOV-2000; 2000US-0246610P.  
 PR 08-NOV-2000; 2000US-0246611P.  
 PR 08-NOV-2000; 2000US-0246613P.  
 PR 17-NOV-2000; 2000US-0249207P.  
 PR 17-NOV-2000; 2000US-0249208P.  
 PR 17-NOV-2000; 2000US-0249209P.  
 PR 17-NOV-2000; 2000US-0249210P.  
 PR 17-NOV-2000; 2000US-0249211P.  
 PR 17-NOV-2000; 2000US-0249212P.  
 PR 17-NOV-2000; 2000US-0249213P.  
 PR 17-NOV-2000; 2000US-0249214P.  
 PR 17-NOV-2000; 2000US-0249215P.  
 PR 17-NOV-2000; 2000US-0249216P.  
 PR 17-NOV-2000; 2000US-0249217P.  
 PR 17-NOV-2000; 2000US-0249218P.  
 PR 17-NOV-2000; 2000US-0249244P.  
 PR 17-NOV-2000; 2000US-0249245P.  
 PR 17-NOV-2000; 2000US-0249246P.  
 PR 17-NOV-2000; 2000US-0249255P.  
 PR 17-NOV-2000; 2000US-0249297P.  
 PR 17-NOV-2000; 2000US-0249299P.  
 PR 17-NOV-2000; 2000US-0249300P.  
 PR 01-DEC-2000; 2000US-0250160P.  
 PR 05-DEC-2000; 2000US-0250391P.  
 PR 05-DEC-2000; 2000US-0251030P.  
 PR 05-DEC-2000; 2000US-0251988P.  
 PR 05-DEC-2000; 2000US-0256719P.  
 PR 06-DEC-2000; 2000US-0251479P.  
 PR 08-DEC-2000; 2000US-0251856P.  
 PR 08-DEC-2000; 2000US-0251868P.  
 PR 08-DEC-2000; 2000US-0251869P.  
 PR 08-DEC-2000; 2000US-0251989P.  
 PR 08-DEC-2000; 2000US-0251990P.  
 PR 11-DEC-2000; 2000US-0254097P.  
 PR 05-JAN-2001; 2001US-0259678P.  
 PR 17-JAN-2001; 2001US-00764860.  
 PR 14-FEB-2002; 2002US-00074095.  
 PA (HUMA-) HUMAN GENOME SCI INC.  
 XX  
 PI Rosen CA, Ruben SM, Barash SC;  
 XX  
 DR WPI; 2003-902033/82.  
 DR N-PSDB; ADG40829.  
 XX  
 PT Novel respiratory system antigen and polynucleotides encoding the  
 PT polypeptides, useful for treating diagnosing, treating or preventing  
 PT tonsillitis, pneumonia, asthma and cystic fibrosis, emphysema, throat  
 PT cancer.  
 XX  
 PS Claim 11; SEQ ID NO 359; 236pp; English.

XX  
 CC The invention describes an isolated polypeptide (I) comprising an amino  
 CC acid sequence that is at least 90% identical to polypeptide fragment of  
 CC any one of 299 respiratory system antigen sequences (PS) and having  
 CC biological activity, polypeptide domain or epitope of PS, full-length  
 CC protein of PS, or variant, allelic variant or species homolog of PS. (I)  
 CC or a polynucleotide (II) encoding (I) is also useful for diagnosing a  
 CC pathological condition or a susceptibility to a pathological condition in  
 CC a subject which involves determining the presence or absence of mutation  
 CC in (II) or determining the presence or amount of expression of (I) in a  
 CC biological sample and diagnosing a pathological condition based on the  
 CC result. The human respiratory system associated polynucleotides, the  
 CC polypeptides encoded by them, and antibodies that immunospecifically bind  
 CC these polypeptides are useful in diagnosis, treatment, prevention and/or  
 CC prognosis of disorders of respiratory system such as throat disorders  
 CC (e.g., vocal cord paralysis, tonsillitis, and laryngitis), lung disorders  
 CC (e.g., pneumonia), allergic disorders, (e.g., asthma and eosinophilic  
 CC pneumonia), pleurisy, cystic fibrosis, emphysema, histiocytosis,  
 CC sarcoidosis, nose disorders (rhinitis and sinusitis), neoplasms and/or  
 CC cancers of respiratory tissues (e.g., throat cancer, lung cancer, and  
 CC cancer of the nose). The polynucleotides are useful in gene therapy

Query Match 32.0%; Score 31; DB 7; Length 50;  
 Best Local Similarity 60.0%; Pred. No. 9.9e+02;  
 Matches 6; Conservative 1; Mismatches 3; Indels 0; Gaps 0;

Qy 2 PNHLSKIAP 11  
 | | | | |  
 Db 36 PPHVNWKTAF 45

RESULT 66  
 ABP71110  
 ID ABP71110 standard; peptide; 19 AA.  
 XX  
 AC ABP71110;  
 XX  
 XX 14-APR-2003 (first entry)  
 DT  
 DE E10 protein CARD region fragment.  
 XX  
 KW BTF3; cell death; apoptosis; basic transcription factor; cytostatic;  
 KW neurotropic; neuroprotective; antiparkinsonian; antiarteriosclerotic;  
 KW antirheumatic; antiarthritic; gene therapy; CARD; E10.  
 XX  
 OS Unidentified.  
 XX  
 PN WO200295001-A2.  
 XX  
 PD 28-NOV-2002.  
 XX  
 PF 21-MAY-2002; 2002MO-US016230.  
 XX  
 PR 21-MAY-2001; 2001US-0292559P.  
 XX  
 XX (REGC ) UNIV CALIFORNIA.  
 PA  
 XX Rothman JH, Bloss T, Witze E;  
 PI WPI; 2003-167228/16.  
 DR  
 XX  
 XX Inhibiting or increasing programmed cell death of a cell, for treating  
 PT e.g. cancer, comprises upregulating or inhibiting, respectively, the  
 PT expression or activity of basic transcription factor (BTF)3 or its  
 PT homolog in the cell.  
 XX  
 XX Example; Fig 2A; 84pp; English.  
 PS  
 CC The invention relates to inhibiting or increasing programmed cell death  
 CC of a cell. The method involves upregulating or inhibiting, respectively,  
 CC the expression or activity of basic transcription factor (BTF)3 or its  
 CC homologue in the cell. The BTF3 polypeptides and nucleic acids are useful  
 CC for inhibiting or increasing programmed cell death. They are used for



CC novel peptides thus avoid potential ovarian damage caused by some  
 CC peptides used as vaccines. The peptides are also useful in assays for  
 CC detecting autoimmune antibodies or for generating antibodies for passive  
 CC immunisation

SQ Sequence 20 AA;  
 Query Match 30.9%; Score 30; DB 2; Length 20;  
 Best Local Similarity 45.5%; Pred. No. 4.9e+02;  
 Matches 5; Conservative 2; Mismatches 4; Indels 0; Gaps 0;

QY 1 EPHNLNSKIAP 11  
 :||| :||| :|||  
 Db 8 DPNELNXCASF 18

RESULT 69  
 ABG76721  
 ID ABG76721 standard; peptide; 24 AA.  
 AC ABG76721;  
 DT 05-NOV-2002 (first entry)  
 XX Hepatitis C virus (HCV)-specific ligand #52.  
 DE Binding specificity; anti-antigen antibody; serum; ADAM;  
 XX antigen detection by antigen mimic; Hepatitis C virus infection; HCV;  
 KW HCV-specific ligand; human.  
 XX Homo sapiens.

OS WO200237115-A1.  
 XX 10-MAY-2002.  
 PN 03-NOV-2000; 2000WO-IT000442.  
 PD 03-NOV-2000; 2000WO-IT000442.  
 XX 03-NOV-2000; 2000WO-IT000442.  
 PR (KENT-) KENTON SRL.  
 XX Felici F, Gargano N, Minenkova O, Monaci P;  
 PI WPI; 2002-599299/64.

DR Making diagnosis of antigen, by identifying binding specificity of anti-  
 PT antigen antibody molecules in serum by antibody detection by antigen  
 PT mimics methodology, and identifying antibodies associated with antigen.  
 XX Claim 26; Page 43; 86pp; English.

CC The present invention relates to a method for making a diagnosis of an  
 CC antigen. The method involves identifying the binding specificity of the  
 CC anti-antigen antibody molecules in serum by the antibody detection by  
 CC antigen mimics (ADAM) methodology. The method comprises screening phage  
 CC libraries using sera from antigen-infected and non-infected individuals,  
 CC and identifying peptides binding antibodies (ligands) specifically  
 CC associated with the antigen. The method of the invention can be used for  
 CC the detection of infectious agents, particularly Hepatitis C virus (HCV).  
 CC The invention provides HCV-specific ligands which are useful for the  
 CC preparation of a diagnostic assay for detecting HCV infection in a  
 CC subject. The HCV-specific ligands are also useful for the preparation of  
 CC vaccines against HCV. ABG76670-ABG76743 represent HCV-specific ligands  
 CC identified from human sera

SQ Sequence 24 AA;  
 Query Match 30.9%; Score 30; DB 5; Length 24;  
 Best Local Similarity 53.8%; Pred. No. 6.1e+02;  
 Matches 7; Conservative 2; Mismatches 4; Indels 0; Gaps 0;

QY 7 SKIAFKIVSQEPA 19

Db 11 SNKAFGIETODPA 23

RESULT 70  
 ABG76719  
 ID ABG76719 standard; peptide; 24 AA.  
 AC ABG76719;  
 XX 05-NOV-2002 (first entry)  
 DT Hepatitis C virus (HCV)-specific ligand #50.

DE Binding specificity; anti-antigen antibody; serum; ADAM;  
 XX antigen detection by antigen mimic; Hepatitis C virus infection; HCV;  
 KW HCV-specific ligand; human.  
 XX Homo sapiens.

OS WO200237115-A1.  
 XX 10-MAY-2002.  
 PN 03-NOV-2000; 2000WO-IT000442.  
 XX 03-NOV-2000; 2000WO-IT000442.  
 PR (KENT-) KENTON SRL.  
 XX Felici F, Gargano N, Minenkova O, Monaci P;  
 PI WPI; 2002-599299/64.

DR Making diagnosis of antigen, by identifying binding specificity of anti-  
 PT antigen antibody molecules in serum by antibody detection by antigen  
 PT mimics methodology, and identifying antibodies associated with antigen.  
 XX Claim 26; Page 43; 86pp; English.

CC The present invention relates to a method for making a diagnosis of an  
 CC antigen. The method involves identifying the binding specificity of the  
 CC anti-antigen antibody molecules in serum by the antibody detection by  
 CC antigen mimics (ADAM) methodology. The method comprises screening phage  
 CC libraries using sera from antigen-infected and non-infected individuals,  
 CC and identifying peptides binding antibodies (ligands) specifically  
 CC associated with the antigen. The method of the invention can be used for  
 CC the detection of infectious agents, particularly Hepatitis C virus (HCV).  
 CC The invention provides HCV-specific ligands which are useful for the  
 CC preparation of a diagnostic assay for detecting HCV infection in a  
 CC subject. The HCV-specific ligands are also useful for the preparation of  
 CC vaccines against HCV. ABG76670-ABG76743 represent HCV-specific ligands  
 CC identified from human sera

SQ Sequence 24 AA;

Query Match 30.9%; Score 30; DB 5; Length 24;  
 Best Local Similarity 47.1%; Pred. No. 6.1e+02;  
 Matches 8; Conservative 2; Mismatches 5; Indels 2; Gaps 1;

QY 3 NHLNSKIAPKIVSQEPA 19  
 :||| :||| :||| :|||  
 Db 9 NYLNK--AFGIEGMOPA 23

RESULT 71  
 AAM16525  
 ID AAM16525 standard; protein; 26 AA.  
 XX AAM16525;  
 AC AAM16525;  
 XX 12-OCT-2001 (first entry)  
 DT

```
DE Peptide #2959 encoded by probe for measuring cervical gene expression.
XX
KW Probe; human; microarray; gene expression; cervical epithelial cell;
KW cervical cancer.
XX
OS Homo sapiens.
XX
FN WO200157278-A2.
XX
XX 09-AUG-2001.
PD
XX
XX 30-JAN-2001; 2001WO-US000670.
PF
XX
XX 04-FEB-2000; 2000US-0180312P.
PR
XX 26-MAY-2000; 2000US-0207456P.
PR
XX 30-JUN-2000; 2000US-00608408.
PR
XX 03-AUG-2000; 2000US-00632366.
PR
XX 21-SEP-2000; 2000US-0234687P.
PR
XX 27-SEP-2000; 2000US-0236359P.
PR
XX 04-OCT-2000; 2000GB-00024263.
XX
XX (MOLE-) MOLECULAR DYNAMICS INC.
PA
XX Penn SG, Hanzel DK, Chen W, Rank DR;
XX
XX WPI; 2001-488901/53.
XX
XX Human genome-derived single exon nucleic acid probes useful for analyzing
XX gene expression in human cervical epithelial cells.
XX
XX Claim 27; SEQ ID NO 21351; 487pp; English.
XX
XX The present invention relates to human single exon nucleic acid probes
XX (SENP; see AAI10068-AA128459). The present sequence is a peptide encoded
XX by one such probe. The SENPs are derived from human HeLa cells. The SENPs
XX can be used to produce a single exon microarray, which can be used for
XX measuring human gene expression in a sample derived from human cervical
XX epithelial cells. By measuring gene expression, the probes are therefore
XX useful in grading and/or staging of diseases of the cervix, notably
XX cervical cancer. Note: The sequence data for this patent did not form
XX part of the printed specification, but was obtained in electronic format
XX directly from WIPO at ftp.wipo.int/pub/published_pct_sequences
XX
XX Sequence 26 AA;
SQ
Query Match 30.9%; Score 30; DB 4; Length 26;
Best Local Similarity 46.2%; Pred. No. 6.7e+02;
Matches 6; Conservative 2; Mismatches 5; Indels 0; Gaps 0;
QY 1 EPNHLSKIAPKI 13
DB 1 EKHLDQMIKFSI 13
XX
RESULT 72
AA04245
ID AA04245 standard; protein; 26 AA.
AC
XX
XX AA04245;
XX
XX 09-OCT-2001 (first entry)
DT
XX
XX Peptide #2927 encoded by probe for measuring breast gene expression.
XX
XX Probe; human; breast disease; breast cancer; development disorder;
KW inflammatory disease; proliferative breast disease; non-carcinoma tumour.
XX
XX Homo sapiens.
OS
XX WO200157270-A2.
FN
XX
XX 09-AUG-2001.
PD
XX
XX
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PF 29-JAN-2001; 2001WO-US000661.
XX
XX 04-FEB-2000; 2000US-0180312P.
PR
XX 26-MAY-2000; 2000US-0207456P.
PR
XX 30-JUN-2000; 2000US-00608408.
PR
XX 03-AUG-2000; 2000US-00632366.
PR
XX 21-SEP-2000; 2000US-0234687P.
PR
XX 27-SEP-2000; 2000US-0236359P.
PR
XX 04-OCT-2000; 2000GB-00024263.
XX
XX (MOLE-) MOLECULAR DYNAMICS INC.
PA
XX Penn SG, Hanzel DK, Chen W, Rank DR;
XX
XX WPI; 2001-476286/51.
XX
XX Novel single exon nucleic acid probe used to measuring gene expression in
XX a human breast.
XX
XX Claim 27; SEQ ID NO 12985; 322pp; English.
XX
XX The present invention relates to novel single exon nucleic acid probes
XX (see AAI00010-AA110067). The present sequence is a peptide encoded by one
XX such probe. The probes are useful for measuring human gene expression in
XX a human breast sample, where the probe hybridises at high stringency to a
XX nucleic acid expressed in the human breast. The probes are useful for
XX predicting diagnosing, grading, staging, monitoring and prognosing
XX diseases of the human breast, particularly those diseases with polygenic
XX aetiology. The diseases include: breast cancer, disorders of development,
XX inflammatory diseases of the breast, fibrocystic changes, proliferative
XX breast disease and non-carcinoma tumours. Note: The sequence data for
XX this patent did not form part of the printed specification, but was
XX obtained in electronic format directly from WIPO at
XX ftp.wipo.int/pub/published_pct_sequences
XX
XX Sequence 26 AA;
SQ
Query Match 30.9%; Score 30; DB 4; Length 26;
Best Local Similarity 46.2%; Pred. No. 6.7e+02;
Matches 6; Conservative 2; Mismatches 5; Indels 0; Gaps 0;
QY 1 EPNHLSKIAPKI 13
DB 1 EKHLDQMIKFSI 13
XX
RESULT 73
AAW8880
ID AAW8880 standard; protein; 27 AA.
AC
XX
XX AAW8880;
XX
XX 01-MAR-1999 (first entry)
DT
XX
XX Polypeptide fragment encoded by gene 100.
DE
XX
XX Human; secreted protein; fusion protein; gene therapy; protein therapy;
KW diagnosis; tissue; cancer; tumour; neurodegenerative disorder; leukaemia;
KW developmental abnormality; foetal deficiency; blood; allergy; renal;
KW immune system; asthma; lymphocytic disease; brain; hepatic; lymphoma;
KW inflammation; ischaemic shock; Alzheimer's disease; restenosis; AIDS;
KW cognitive disorder; schizophrenia; prostate; obesity; osteoclast; thymus;
KW osteoporosis; arthritis; testis; lung; thyroiditis; thyroid; digestion;
KW endocrine; metabolism; regulation; malabsorption; gastritis; neoplasm.
XX
XX Homo sapiens.
OS
XX WO9854963-A2.
FN
XX
XX 10-DEC-1998.
PD
XX
XX 04-JUN-1998; 98WO-US011422.
PF
XX
```



```
XX OS Homo sapiens.
XX PN WO3854963-A2.
XX PD 10-DEC-1998.
XX PF 04-JUN-1998; 98WO-US011422.
XX PR 06-JUN-1997; 97US-0048875P.
XX PR 06-JUN-1997; 97US-0048876P.
XX PR 06-JUN-1997; 97US-0048877P.
XX PR 06-JUN-1997; 97US-0048878P.
XX PR 06-JUN-1997; 97US-0048880P.
XX PR 06-JUN-1997; 97US-0048881P.
XX PR 06-JUN-1997; 97US-0048882P.
XX PR 06-JUN-1997; 97US-0048883P.
XX PR 06-JUN-1997; 97US-0048884P.
XX PR 06-JUN-1997; 97US-0048885P.
XX PR 06-JUN-1997; 97US-0048882P.
XX PR 06-JUN-1997; 97US-0048893P.
XX PR 06-JUN-1997; 97US-0048894P.
XX PR 06-JUN-1997; 97US-0048895P.
XX PR 06-JUN-1997; 97US-0048896P.
XX PR 06-JUN-1997; 97US-0048898P.
XX PR 06-JUN-1997; 97US-0048899P.
XX PR 06-JUN-1997; 97US-0048900P.
XX PR 06-JUN-1997; 97US-0048901P.
XX PR 06-JUN-1997; 97US-0048915P.
XX PR 06-JUN-1997; 97US-0048916P.
XX PR 06-JUN-1997; 97US-0048917P.
XX PR 06-JUN-1997; 97US-0048919P.
XX PR 06-JUN-1997; 97US-0048949P.
XX PR 06-JUN-1997; 97US-0048962P.
XX PR 06-JUN-1997; 97US-0048963P.
XX PR 06-JUN-1997; 97US-0048970P.
XX PR 06-JUN-1997; 97US-0048971P.
XX PR 06-JUN-1997; 97US-0048972P.
XX PR 06-JUN-1997; 97US-0048974P.
XX PR 06-JUN-1997; 97US-0049019P.
XX PR 06-JUN-1997; 97US-0049020P.
XX PR 06-JUN-1997; 97US-0049373P.
XX PR 06-JUN-1997; 97US-0049374P.
XX PR 06-JUN-1997; 97US-0049375P.
XX PR 05-SEP-1997; 97US-0057584P.
XX PR 05-SEP-1997; 97US-0057627P.
XX PR 05-SEP-1997; 97US-0057628P.
XX PR 05-SEP-1997; 97US-0057629P.
XX PR 05-SEP-1997; 97US-0057634P.
XX PR 05-SEP-1997; 97US-0057635P.
XX PR 05-SEP-1997; 97US-0057642P.
XX PR 05-SEP-1997; 97US-0057643P.
XX PR 05-SEP-1997; 97US-0057644P.
XX PR 05-SEP-1997; 97US-0057645P.
XX PR 05-SEP-1997; 97US-0057646P.
XX PR 05-SEP-1997; 97US-0057647P.
XX PR 05-SEP-1997; 97US-0057648P.
XX PR 05-SEP-1997; 97US-0057649P.
XX PR 05-SEP-1997; 97US-0057650P.
XX PR 05-SEP-1997; 97US-0057651P.
XX PR 05-SEP-1997; 97US-0057654P.
XX PR 05-SEP-1997; 97US-0057661P.
XX PR 05-SEP-1997; 97US-0057662P.
XX PR 05-SEP-1997; 97US-0057666P.
XX PR 05-SEP-1997; 97US-0057667P.
XX PR 05-SEP-1997; 97US-0057668P.
XX PR 05-SEP-1997; 97US-0057760P.
XX PR 05-SEP-1997; 97US-0057761P.
XX PR 05-SEP-1997; 97US-0057762P.
XX PR 05-SEP-1997; 97US-0057763P.
XX PR 05-SEP-1997; 97US-0057764P.
XX PR 05-SEP-1997; 97US-0057765P.

PR 05-SEP-1997; 97US-0057769P.
PR 05-SEP-1997; 97US-0057770P.
PR 05-SEP-1997; 97US-0057771P.
PR 05-SEP-1997; 97US-0057774P.
PR 05-SEP-1997; 97US-0057775P.
PR 05-SEP-1997; 97US-0057776P.
PR 05-SEP-1997; 97US-0057777P.
PR 05-SEP-1997; 97US-0057778P.
PR 18-DEC-1997; 97US-0070923P.
XX (HUMA-) HUMAN GENOME SCI INC.
XX Young P, Greene JM, Perrie AM, Ruben SM, Rosen CA, Hu J;
XX Olsen HS, Ebner R, Brewer LA, Moore PA, Shi Y, Florence C;
XX Florence K, Lafleur DW, Ni J, Fan P, Wei Y, Fischer CL, Soppet DR;
XX Li Y, Zeng Z, Kyaw H, Yu G, Feng P, Dillon PL, Endress GA;
XX Carter KC;
XX WPI; 1999-059865/05.
XX N-PSDB; AAV84510.
XX New isolated human genes and the secreted polypeptides they encode -
XX useful for diagnosis and treatment of e.g. cancers, neurological
XX disorders, immune diseases, inflammation or blood disorders.
XX Disclosure; Page 77; 772pp; English.
XX The invention relates to nucleic acid sequences (AAV84411 to AAV84633)
XX encoding human secreted proteins (AAW88534 to AAW88756). The secreted
XX protein gene sequences are deposited with the ATCC under deposit numbers
XX ATCC 97979, 97974, 97975, 97977, 209007, 209008, 209009, 209010,
XX 209011, 209080, 209081, 209082, 209083, 209084, 209085, 209511. Host
XX cells comprising recombinant vectors containing the nucleic acid
XX sequences are used for the recombinant production of the secreted
XX proteins. The polynucleotide and amino acid sequences are useful for are
XX useful for preventing, treating or ameliorating medical conditions e.g.
XX by protein or gene therapy. Pathological conditions can be also diagnosed
XX by determining the amount of the new polypeptides in a sample or by
XX determining the presence of mutations in the new polynucleotides.
XX Specific uses are described for each of the polynucleotides, based on
XX which tissues they are most highly expressed in, and include developing
XX products for the diagnosis or treatment of cancer, neurodegenerative
XX disorders, developmental abnormalities and foetal deficiencies, blood
XX disorders, tumours, leukemias, diseases of the immune system, autoimmune
XX diseases, hepatic and renal disease, lymphomas, inflammation, allergies,
XX ischemic shock, Alzheimer's and cognitive disorders, schizophrenia,
XX retinosis, prostate diseases, obesity, disorders involving osteoclasts
XX such as osteoporosis, arthritis or malignancies, diseases of testes, lung
XX or thymus, digestive/endocrine disorders, infections and AIDS. The
XX polypeptides are also useful for identifying their binding partners. The
XX present sequence represents a polypeptide fragment encoded by a gene of
XX the invention (see descriptor line for gene number)
XX Sequence 27 AA;
XX Query Match 30.9%; Score 30; DB 2; Length 27;
XX Best Local Similarity 37.5%; Pred. No. 7.1e+02;
XX Matches 6; Conservative 4; Mismatches 6; Indels 0; Gaps 0;
QY 2 PNHLSKIAFKIVSQE 17
Db 6 PSANNQRFAPSLSEE 21
RESULT 75
ABB50945
ID ABB50945 standard; protein; 27 AA.
XX
XX ABB50945;
XX
XX 07-FEB-2002 (first entry)
XX Human secreted protein encoded by gene 100 SEQ ID NO:898.
DE
```







Sequence 27 AA;



Sequence 27 AA;

QY	30.9%; Score 30; DB 7; Length 27; Best Local Similarity 37.5%; Pred. No. 7.1e+02; Matches 6; Conservative 4; Mismatches 6; Indels 0; Gaps 0;
DB	2 PNHLNSKIAFKIVSOE 17 6 PSANNQRFAPSLSEE 21
RESULT 81	
ID	ABP26325
AC	ABP26325 standard; protein; 30 AA.
XX	ABP26325;
DT	02-JUL-2002 (first entry)
DE	Streptococcus polypeptide SEQ ID NO 1826.
XX	Streptococcus; GAS; GBS; group B streptococcus; Streptococcus agalactiae;
KW	group A streptococcus; Streptococcus pyogenes; antibacterial;
KW	antiinflammatory; infection; vaccine; meningitis; gene therapy.
XX	Streptococcus agalactiae.
OS	
PN	WO200234771-A2.
XX	
PD	02-MAY-2002.
XX	
PF	29-OCT-2001; 2001WO-GB004789.
XX	
FR	27-OCT-2000; 2000GB-00026333.
PR	24-NOV-2000; 2000GB-00028727.
PR	07-MAR-2001; 2001GB-00005640.
XX	(CHIR-) CHIRON SPA.
PA	(GENO-) INST GENOMIC RES.
XX	
PI	Telford J, Masignani V, Margarit Y RosI, Grandi G, Fraser C;
PI	Tetelin H;
XX	
DR	WPI; 2002-352536/38.
DR	N-PSDB; ABN66956.
XX	
PT	New Streptococcus protein for the treatment or prevention of infection or
PT	disease caused by Streptococcus bacteria, such as meningitis, and for
PT	detecting a compound that binds to the protein.
XX	
PS	Claim 1; Page 3332; 4525pp; English.
XX	
CC	The invention relates to a protein (ABP25413-ABP30895) from group B
CC	streptococcus/GBS (Streptococcus agalactiae) or group A streptococcus/GAS
CC	(Streptococcus pyogenes), comprising one of 5483 sequences (S1), given in
CC	the specification. The proteins have antibacterial and antiinflammatory
CC	activity. (I), nucleic acids encoding (I), ABN66044-ABN71526 and
CC	antibodies that bind (I) are used in the manufacture of medicaments for
CC	the treatment or prevention of infection or disease caused by
CC	Streptococcus bacteria, particularly S. agalactiae and S. pyogenes.
CC	Nucleic acids encoding (I) are used to detect Streptococcus in a
CC	biological sample. (I) is used to determine whether a compound binds to
CC	(I). A composition comprising (I) or a nucleic acid encoding (I), may be
CC	used as a vaccine or diagnostic composition. The disease caused by
CC	Streptococcus that is prevented or treated may be meningitis. Nucleic
CC	acid encoding (I) may be used to recombinantly produce (I) and may be
CC	used in gene therapy. Antibodies to (I) are used for affinity
CC	chromatography, immunoassays, and distinguishing/identifying
CC	Streptococcus proteins
XX	
SQ	Sequence 30 AA;

QY	30.9%; Score 30; DB 5; Length 30; Best Local Similarity 66.7%; Pred. No. 8e+02; Matches 6; Conservative 2; Mismatches 1; Indels 0; Gaps 0;
DB	7 SKIAPKIVS 15 4 SKISFKIAT 12
RESULT 82	
ID	AAU84722 standard; peptide; 30 AA.
XX	AAU84722;
DT	08-MAY-2002 (first entry)
XX	
DE	HCV HepC1a segment 125.
XX	
KW	Savine; vaccine; cancer; viral infection; HIV; hepatitis C virus;
KW	viral infection; human immunodeficiency virus; melanoma;
KW	bacterial infection; Salmonella; Legionella; parasitic infection;
KW	Trypanosoma; Toxoplasma; Giardia.
XX	
OS	Hepatitis C virus.
XX	
PN	WO200190197-A1.
XX	
PD	29-NOV-2001.
XX	
PF	25-MAY-2001; 2001WO-AU000622.
XX	
PR	26-MAY-2000; 2000AU-00007761.
XX	(AUSU ) UNIV AUSTRALIAN NAT.
PA	Thomson SA, Ramshaw IA;
PI	
XX	
DR	WPI; 2002-147575/19.
DR	N-PSDB; ABK36560.
XX	
PT	New synthetic polypeptides having several different segments of at least
PT	one parent polypeptide linked together differently compared to the
PT	linkage in the parent polypeptide, for inducing immune response against a
PT	pathogen or cancer.
XX	
PS	Example 2; Fig 26; 364pp; English.
XX	
CC	The invention relates to a new synthetic polypeptide (I) comprising
CC	several different segments of at least one parent polypeptide linked
CC	together in a different relationship relative to their linkage in the
CC	parent polypeptide to impede, abrogate or otherwise alter at least one
CC	function associated with the parent polypeptide and for inducing an
CC	immune response against a pathogen or cancer. Also included are a
CC	synthetic polynucleotide encoding and a computer system for designing the
CC	synthetic polypeptides. The synthetic polypeptides and polynucleotides
CC	are referred to as a Savine. The synthetic polypeptide is useful for a
CC	modulating immune responses preferably directed against a pathogen or a
CC	cancer, (e.g., cancers of the lung, breast, ovary, cervix, colon, head
CC	and neck, pancreas, prostate, stomach, bladder, kidney, bone liver,
CC	oesophagus, brain, testicle, uterus), as potentiating agents.
CC	Compositions comprising the polypeptide may be used in the treatment or
CC	prophylaxis against viral (such as infections caused by HIV (human
CC	immunodeficiency virus), hepatitis, influenza, Japanese encephalitis
CC	virus, Epstein-Barr virus and respiratory syncytial virus), bacterial
CC	(e.g., infections caused by Neisseria, Meningococcal, Haemophilus,
CC	Salmonella, Streptococcal, Legionella and Mycobacterium or parasitic
CC	(e.g., infections caused by Plasmodium, Schistosoma, Leishmania,
CC	Trypanosoma, Toxoplasma and Giardia) infections. The present sequence is
CC	a peptide derived from a parent protein used to construct a savine of the
CC	invention
XX	
SQ	Sequence 30 AA;

QY	30.9%; Score 30; DB 5; Length 30; Best Local Similarity 66.7%; Pred. No. 8e+02; Matches 6; Conservative 2; Mismatches 1; Indels 0; Gaps 0;
DB	7 SKIAPKIVS 15 4 SKISFKIAT 12
RESULT 82	
ID	AAU84722 standard; peptide; 30 AA.
XX	AAU84722;
DT	08-MAY-2002 (first entry)
XX	
DE	HCV HepC1a segment 125.
XX	
KW	Savine; vaccine; cancer; viral infection; HIV; hepatitis C virus;
KW	viral infection; human immunodeficiency virus; melanoma;
KW	bacterial infection; Salmonella; Legionella; parasitic infection;
KW	Trypanosoma; Toxoplasma; Giardia.
XX	
OS	Hepatitis C virus.
XX	
PN	WO200190197-A1.
XX	
PD	29-NOV-2001.
XX	
PF	25-MAY-2001; 2001WO-AU000622.
XX	
PR	26-MAY-2000; 2000AU-00007761.
XX	(AUSU ) UNIV AUSTRALIAN NAT.
PA	Thomson SA, Ramshaw IA;
PI	
XX	
DR	WPI; 2002-147575/19.
DR	N-PSDB; ABK36560.
XX	
PT	New synthetic polypeptides having several different segments of at least
PT	one parent polypeptide linked together differently compared to the
PT	linkage in the parent polypeptide, for inducing immune response against a
PT	pathogen or cancer.
XX	
PS	Example 2; Fig 26; 364pp; English.
XX	
CC	The invention relates to a new synthetic polypeptide (I) comprising
CC	several different segments of at least one parent polypeptide linked
CC	together in a different relationship relative to their linkage in the
CC	parent polypeptide to impede, abrogate or otherwise alter at least one
CC	function associated with the parent polypeptide and for inducing an
CC	immune response against a pathogen or cancer. Also included are a
CC	synthetic polynucleotide encoding and a computer system for designing the
CC	synthetic polypeptides. The synthetic polypeptides and polynucleotides
CC	are referred to as a Savine. The synthetic polypeptide is useful for a
CC	modulating immune responses preferably directed against a pathogen or a
CC	cancer, (e.g., cancers of the lung, breast, ovary, cervix, colon, head
CC	and neck, pancreas, prostate, stomach, bladder, kidney, bone liver,
CC	oesophagus, brain, testicle, uterus), as potentiating agents.
CC	Compositions comprising the polypeptide may be used in the treatment or
CC	prophylaxis against viral (such as infections caused by HIV (human
CC	immunodeficiency virus), hepatitis, influenza, Japanese encephalitis
CC	virus, Epstein-Barr virus and respiratory syncytial virus), bacterial
CC	(e.g., infections caused by Neisseria, Meningococcal, Haemophilus,
CC	Salmonella, Streptococcal, Legionella and Mycobacterium or parasitic
CC	(e.g., infections caused by Plasmodium, Schistosoma, Leishmania,
CC	Trypanosoma, Toxoplasma and Giardia) infections. The present sequence is
CC	a peptide derived from a parent protein used to construct a savine of the
CC	invention
XX	
SQ	Sequence 30 AA;

Matches 6; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

Qy 9 IAFKIVSQE 17  
:||||:|  
Db 6 VAFKIMSGE 14

## RESULT 83

AAU84721  
AAU84721 standard; peptide; 30 AA.

AC AAU84721;

DT 08-MAY-2002 (first entry)

DE HCV HepC1a segment 124.

XX Savine; vaccine; cancer; viral infection; HIV; hepatitis C virus;  
XX viral infection; human immunodeficiency virus; melanoma;  
XX bacterial infection; Salmonella; Legionella; parasitic infection;  
XX Trypanosoma; Toxoplasma; Giardia.

OS Hepatitis C virus.

XX WO200190197-A1.

XX 29-NOV-2001.

XX 25-MAY-2001; 2001WO-AUC00622.

XX 26-MAY-2000; 2000AU-00007761.

XX (AUSU) UNIV AUSTRALIAN NAT.

XX Thomson SA, Ramshaw IA;

XX WPI; 2002-147575/19.

XX N-PSDB; ABK36559.

XX New synthetic polypeptides having several different segments of at least  
PT one parent polypeptide linked together differently compared to the  
PT linkage in the parent polypeptide, for inducing immune response against a  
PT pathogen or cancer.

XX Example 2; Fig 26; 364pp; English.

XX The invention relates to a new synthetic polypeptide (I) comprising  
XX several different segments of at least one parent polypeptide linked  
XX together in a different relationship relative to their linkage in the  
XX parent polypeptide to impede, abrogate or otherwise alter at least one  
XX function associated with the parent polypeptide and for inducing an  
XX immune response against a pathogen or cancer. Also included are a  
XX synthetic polynucleotide encoding and a computer system for designing the  
XX synthetic polypeptides. The synthetic polypeptides and polynucleotides  
XX are referred to as a vaccine. The synthetic polypeptide is useful for  
XX modulating immune responses preferably directed against a pathogen or a  
XX cancer, (e.g., cancers of the lung, breast, ovary, cervix, colon, head  
XX and neck, pancreas, prostate, stomach, bladder, kidney, bone liver,  
XX oesophagus, brain, testicle, uterus), as potentiating agents.  
XX Compositions comprising the polypeptide may be used in the treatment or  
XX prophylaxis against viral (such as infections caused by HIV (human  
XX immunodeficiency virus), hepatitis, influenza, Japanese encephalitis  
XX virus, Epstein-Barr virus and respiratory syncytial virus), bacterial  
XX (e.g., infections caused by Neisseria, Meningococcus, Haemophilus,  
XX Salmonella, Streptococcus, Legionella and Mycobacterium or parasitic  
XX (e.g., infections caused by Plasmodium, Schistosoma, Leishmania,  
XX Trypanosoma, Toxoplasma and Giardia) infections. The present sequence is  
XX a peptide derived from a parent protein used to construct a vaccine of the  
XX invention

XX Sequence 30 AA;

Query Match 30.9%; Score 30; DB 5; Length 30;

Best Local Similarity 66.7%; Pred. No. 8e+02; Indels 0; Gaps 0;  
Matches 6; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

Qy 9 IAFKIVSQE 17  
:||||:|  
Db 21 VAFKIMSGE 29

## RESULT 84

ABG68718  
ID ABG68718 standard; protein; 30 AA.

XX AC ABG68718;

XX 07-OCT-2002 (first entry)

XX Human prostate specific protein DEX0293\_87.

XX Human; prostate specific nucleic acid; PSNA; prostate cancer; PSP;  
XX prostate specific protein; cytostatic; non-cancerous prostate disease;  
XX gene therapy; cancer; immunostimulant; vaccine.

XX Homo sapiens.

XX WO200255735-A2.

XX 18-JUL-2002.

XX 27-NOV-2001; 2001WO-US044363.

XX 27-NOV-2000; 2000US-0253176P.

XX (DIAD-) DIADEXUS INC.

XX Salceda S, Macina RA, Recipon H, Caferkey R, Ali S, Sun Y;

XX Liu C, Chen S;

XX WPI; 2002-557831/59.

XX N-PSDB; ABK97600.

XX New prostate specific genes, useful for treating or diagnosing cancer, or  
PT useful as vaccines for treating cancer, particularly prostate cancer, in  
PT a patient.

XX Claim 11; Page 196-197; 212pp; English.

XX The invention relates to a new isolated prostate-specific nucleic acid  
XX (PSNA) molecule comprising the cDNA sequences appearing as ABK97574-  
XX ABK97642 which encode prostate specific proteins appearing as ABG68701-  
XX ABG68746, or a sequence hybridising to a PSNA or which has 60% sequence  
XX homology with a PSNA. Also included are a method of determining the  
XX presence of a PSNA in a sample, a vector comprising the PSNA, a host cell  
XX comprising the vector, producing the polypeptide encoded by the PSNA, a  
XX method of determining the presence of a PSP in a sample, diagnosing and  
XX monitoring the presence and metastases of prostate cancer in a patient, a  
XX kit for detecting a risk of cancer or presence of cancer in a patient  
XX (the kit comprising a means for determining the presence of the PSNA or  
XX PSP in a sample of a patient) and a vaccine comprising the polypeptide or  
XX the nucleic acid encoding the polypeptide. The PSNA, PSP and anti-PSP  
XX antibody are useful for diagnosing and treating cancer in a patient (e.g.  
XX by gene therapy). The nucleic acid molecule and polypeptide are also  
XX useful as vaccines for treating cancer, particularly prostate cancer and  
XX non-cancerous prostate diseases. The present sequence is a PSP of the  
XX invention

XX Sequence 30 AA;

Query Match 30.9%; Score 30; DB 5; Length 30;

Best Local Similarity 45.5%; Pred. No. 8e+02;  
Matches 5; Conservative 5; Mismatches 1; Indels 0; Gaps 0;

Qy 7 SKIAFKIVSQE 17  
:||||:|





XX OS Homo sapiens.  
 XX PN WO200122920-A2.  
 XX PD 05-APR-2001.  
 XX PF 28-SEP-2000; 2000WO-US026524.  
 XX PR 29-SEP-1999; 99US-0157137P.  
 XX PR 03-NOV-1999; 99US-0163280P.  
 XX PA (HUMA-) HUMAN GENOME SCI INC.  
 XX PI Ruben SM, Barash SC, Birse CE, Rosen CA;  
 XX WPI; 2001-235357/24.  
 XX DR N-PSDB; AAH34599.  
 XX Nucleic acids encoding 4277 human colon cancer-associated polypeptides,  
 PT useful for preventing, diagnosing and/or treating colorectal cancers.  
 PT Claim 11; Page 7465; 9803pp; English.  
 XX AAH32943 to AAH37195 and AAH37788 represent human colon  
 CC cancer-associated nucleic acid molecules (N) and proteins (P), where the  
 CC proteins are collectively known as colon cancer antigens. The colon  
 CC cancer antigens have cytostatic activity and can be used in gene therapy  
 CC and vaccine production. N and P may be used in the prevention, diagnosis  
 CC and treatment of diseases associated with inappropriate P expression. For  
 CC example, N and P may be used to treat disorders associated with decreased  
 CC expression by rectifying mutations or deletions in a patient's genome  
 CC that affect the activity of P by expressing inactive proteins or to  
 CC supplement the patients own production of P. Additionally, N may be used  
 CC to produce the colon cancer-associated Ps, by inserting the nucleic acids  
 CC into a host cell and culturing the cell to express the proteins. N and P  
 CC can be used in the prevention, diagnosis and treatment of colorectal  
 CC carcinomas and cancers. AAH37196 to AAH37204 and AAH37789 represent  
 CC sequences used in the exemplification of the present invention. N.B.  
 CC Pages 666 to 682 and page 7053 of the sequence listing were missing at  
 CC time of publication, meaning no sequences are present for SEQ ID NO:1027  
 CC to 1052, 7921 and 7922  
 XX Sequence 32 AA;  
 SQ  
 Query Match 30.9%; Score 30; DB 4; Length 32;  
 Best Local Similarity 77.8%; Pred.No. 8.7e+02;  
 Matches 7; Conservative 1; Mismatches 1; Indels 0; Gaps 0;  
 QY 8 KIAFKIVSQ 16  
 Db |||:||||  
 5 KIAWKIVQ 13  
 RESULT 88  
 ADH96838  
 ID ADH96838 standard; protein; 32 AA.  
 XX AC ADH96838;  
 XX DT 06-MAY-2004 (first entry)  
 XX DE S. pneumoniae DNA polymerase III beta-subunit epitope #2.  
 XX antibacterial; antiinflammatory; gastrointestinal; antiulcer;  
 KW antiarrheic; ophthalmological; enzyme inhibitor; antisense therapy;  
 KW vaccine; microbial target; modulator; furuncle; pneumonia; gastritis;  
 KW peptic ulcer disease; diarrhoea; meningitis; bacteraemia; conjunctivitis;  
 KW toxic shock syndrome; epitope.  
 XX Streptococcus pneumoniae.  
 OS WO2003102190-A2.  
 PN

XX PD 11-DEC-2003.  
 XX PF 02-JUN-2003; 2003WO-CA000786.  
 XX PR 31-MAY-2002; 2002US-0384634P.  
 XX PR 31-MAY-2002; 2002US-0385157P.  
 XX PR 04-JUN-2002; 2002US-0385542P.  
 XX PR 04-JUN-2002; 2002US-0385611P.  
 XX PR 04-JUN-2002; 2002US-0385747P.  
 XX PR 04-JUN-2002; 2002US-0385750P.  
 XX PR 04-JUN-2002; 2002US-0385752P.  
 XX PR 04-JUN-2002; 2002US-0385773P.  
 XX PR 04-JUN-2002; 2002US-0385780P.  
 XX PR 04-JUN-2002; 2002US-0385785P.  
 XX PR 04-JUN-2002; 2002US-0385797P.  
 XX PR 05-JUN-2002; 2002US-0386022P.  
 XX PR 05-JUN-2002; 2002US-0386024P.  
 XX PR 05-JUN-2002; 2002US-0386087P.  
 XX PR 05-JUN-2002; 2002US-0386141P.  
 XX PR 05-JUN-2002; 2002US-0386350P.  
 XX PR 05-JUN-2002; 2002US-0386586P.  
 XX PR 06-JUN-2002; 2002US-0386368P.  
 XX PR 06-JUN-2002; 2002US-0386389P.  
 XX PR 06-JUN-2002; 2002US-0386436P.  
 XX PR 06-JUN-2002; 2002US-0386441P.  
 XX PR 06-JUN-2002; 2002US-0386528P.  
 XX PR 06-JUN-2002; 2002US-0386573P.  
 XX PR 06-JUN-2002; 2002US-0386834P.  
 XX PR 31-JUL-2002; 2002US-0399839P.  
 XX PR 31-JUL-2002; 2002US-0399861P.  
 XX PR 31-JUL-2002; 2002US-0399969P.  
 XX PR 31-JUL-2002; 2002US-0399970P.  
 XX PR 31-JUL-2002; 2002US-0399983P.  
 XX PR 31-JUL-2002; 2002US-0399984P.  
 XX PR 31-JUL-2002; 2002US-0399985P.  
 XX PR 01-AUG-2002; 2002US-0400154P.  
 XX PR 01-AUG-2002; 2002US-0400230P.  
 XX PR 01-AUG-2002; 2002US-0400268P.  
 XX PR 01-AUG-2002; 2002US-0400363P.  
 XX PR 01-AUG-2002; 2002US-0400365P.  
 XX PR 01-AUG-2002; 2002US-0400374P.  
 XX PR 01-AUG-2002; 2002US-0400380P.  
 XX PR 01-AUG-2002; 2002US-0400433P.  
 XX PR 01-AUG-2002; 2002US-0400434P.  
 XX PR 01-AUG-2002; 2002US-0400436P.  
 XX PR 01-AUG-2002; 2002US-0400442P.  
 XX PR 01-AUG-2002; 2002US-0400463P.  
 XX PA (AFFI-) AFFINIUM PHARM INC.  
 XX Edwards A, Dharamsi A, Vedadi M, Vallee F, Awrey D, Beattie B;  
 PI Richards D, Domagala M, Mansoury K, Virag C, Buzadziya K;  
 PI McDonald M, Houston S, Arrowsmith C, Ouyang H, Nethery K, Ng I;  
 PI Kanagarajah D;  
 XX WPI; 2004-071165/07.  
 XX Compositions comprising recombinant polypeptide targets for pathogenic  
 PT bacteria, useful for designing modulators for preventing or treating a  
 PT disease or disorder associated with the species of origin for the  
 PT polypeptide.  
 XX Disclosure; SEQ ID NO 29; 606pp; English.  
 XX The invention relates to novel compositions (I) comprising isolated,  
 XX recombinant polypeptides, amino acid sequences having at least about 95%  
 CC identity with these or an amino acid sequence encoded by a polynucleotide  
 CC that hybridizes under stringent conditions to the complementary strand of  
 CC the polynucleotide encoding these polypeptides. The compositions and  
 CC polypeptides are useful as microbial targets for designing modulators for  
 CC the prevention or treatment of a disease or disorder associated with the

species of origin for the polypeptide, e.g. furuncle, pneumonia, gastritis, peptic ulcer disease, diarrhoea, meningitis, bacteraemia, conjunctivitis or toxic shock syndrome. The polypeptides are also useful for diagnosing a patient suffering from a disease or disorder of a pathogenic species, or for monitoring the effectiveness of an antipathogenic treatment. This sequence corresponds to an epitope of one of the protein sequences of the invention.

Sequence 32 AA;

Query Match	30.9%;	Score 30;	DB 8;	Length 32;
Best Local Similarity	45.5%;	Pred. No. 8.7e+02;		
Matches 5;	Conservative 3;	Mismatches 3;	Indels 0;	Gaps 0;

9 IAFKIVSOEPA 19

14 VDFKVFSEAPA 24

RESULT 89  
AAW29908  
ID AAW29908 standard: peptide: 33 AA:

AC AAW29908:

16-MAR-1998 (first entry)

DE porcine AMPK- $\alpha$  active peptide fragment 3.

5'-AMP activated protein kinase; AMPK; catalytic subunit;  
protein phosphorylation; cholesterol; fatty acid; pig;  
hormone sensitive lipase; HSL; alpha subunit

OS Sus scrofa.

PN W09725341-A1.

17-JUL-1997

07-JAN-1997: 97WO-US0000270.

PR 08-JAN-1996; 96AU-00007450.

PA (SVIN-) ST VINCENTS INST MEDICAL RES.  
PA (DART-) DARTMOUTH COLLEGE.

PT Kemp BE. Stapleton DI. Mitchelhill KI. Witters LA;

WPT: 1997-372811/34.

New isolated 5'-AMP-activated protein kinase subunit(s) - used to develop products for treating e.g. hyper-cholesterolaemia, obesity, hypoxia, ischaemia, nutrition disorders or diabetes mellitus.

ps Disclosure: Page 36: 63pp: English.

This sequence represent a biologically active peptide derived from the 5'-AMP-activated protein kinase (AMPK) catalytic alpha subunit from pig liver. This fragment retains at least one of the activities of native AMPK-alpha i.e the ability to stimulate phosphorylation of protein molecules and the ability to mimic the binding of native AMPK-alpha to at least one antibody or ligand molecule. AMPK polypeptides can be used to identify compounds which regulate the action of kinases. Such fragments can be used to reduce biosynthesis of cholesterol and fatty acids. They may also be used to inhibit the release of these molecules from intracellular stores by hormone sensitive lipase (HSL). They may also be used to reduce cellular malonyl CoA levels and promote the beta-oxidation of fatty acids by mitochondria. AMPK-alpha fragments could be used in the treatment of e.g. hypercholesterolaemia, hyperlipidaemia, obesity, clinical syndromes associated with hypoxia or ischaemia (e.g. myocardial infarction) disorders of nutrition and diabetes mellitus

Sequence 33 AA:

QY	4	HLNSKIA	10
		:   :	
pb	12	HMNAKIA	18

RESULT 90  
AAW35030  
IN AAW35030 standard: peptide: 33 AA.

AC AAW35030;

DT 22-MAY-1998 (first entry)

Water soluble beta-sheet forming peptide beta pep-13.

water soluble; beta-sheet; treatment; acute peritonitis; fever; shock;  
bacterial infection: tumour.

synthetic.

PN WO9744354-A2.

27-NOV-1997.

23-MAY-1997:

24-MAY-1996: 96US-00653632.

XX  
EK

[illegible]

Pt Gray Bk., Haslemar Cr., Mayo Rf.  
XX

XX  
XX

UNCLAS//NOFORN

PT endotoxin, acts as bactericidal agent and inhibits TNF-alpha levels and endothelial cell proliferation.

xx ps Claim 21: page 22: 60pp: English.

Synthesising a water soluble peptide, e.g. the present peptide, where at least 35% of its amino acids have hydrophobic side chains comprises combining amino acids with non-charged polar and charged side chains, where the ratio of positively to negatively charged side chains is at least about 2:1, with amino acids having hydrophobic side chains. A peptide prepared using the above process is capable of neutralising endotoxin, and is active as bactericidal agent. It may also be used to inhibit tumour necrosis factor- $\alpha$  (TNF- $\alpha$ ) levels and endothelial cell proliferation and promote intercellular adhesion molecule (ICAM) expression. The peptide may therefore be used to treat acute peritonitis, fever, shock and bacterial infections. It may also be used to inhibit angiogenesis, e.g. in treatment of tumours. The peptide is water soluble, soluble under physiological conditions, forms beta-sheets and can self associate.

Sequence 33 AA:

Query Match 30.9%; Score 30; DB 2; Length 33;  
Best Local Similarity 60.0%; Pred. No. 9e+02;  
Matches 6: Conservative 1: Mismatches 3; Indels

0v 4 HLNSKIAFKI 13

db	15	HLKWKIIFKL	24
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## RESULT 91

```

ABM73359
ID ABM73359 standard; protein; 34 AA.
XX
AC ABM73359;
XX
DT 20-NOV-2003 (first entry)
DE
DE Staphylococcus aureus protein #2599.
XX
KW Antibacterial; vaccine; gene therapy; infection; sepsis; diagnosis;
KW enzymatic assay; antibiotic target.
XX
OS Staphylococcus aureus.
XX
PN WO200294868-A2.
XX
PD 28-NOV-2002.
XX
XX 27-MAR-2002; 2002WO-IB002637.
FF
XX 27-MAR-2001; 2001GB-00007661.
XX
PA (CHIR-) CHIRON SPA.
XX
PI Masignani V, Mora M, Scarselli M;
XX
DR WPI; 2003-120786/11.
DR N-PSDB; ACF74919.
XX
XX New Staphylococcus aureus protein, useful as a vaccine for treating or
PT preventing Staphylococcal infection, specifically an infection caused by
PT S. aureus, e.g. sepsis.
XX
XX Claim 1; SEQ ID NO 5198; 49bp; English.
XX
XX The invention relates to novel genes and encoded proteins from
CC Staphylococcus aureus. A composition comprising the S. aureus protein, a
CC nucleic acid encoding the protein, or an antibody to the protein, is
CC useful as a pharmaceutical, particularly as a vaccine for treating or
CC preventing infection due to Staphylococcus bacteria, specifically an
CC infection caused by S. aureus. The composition is particularly useful for
CC treating or preventing sepsis in a patient. The composition can also be
CC used for diagnostics. The protein is also used in an assay for enzymatic
CC studies and as a target for antibiotics. This sequence represents one of
CC the novel S. aureus proteins of the invention
XX
SQ Sequence 34 AA;
Query Match 30.9%; Score 30; DB 6; Length 34;
Best Local Similarity 46.7%; Pred. No. 9.4e+02;
Matches 7; Conservative 1; Mismatches 7; Indels 0; Gaps 0;

QY 2 PNHLSKIAPKIVSQ 16
||| |||
Db 15 PNIKTRKAUKIKQ 29

RESULT 92
ABM74126
ID ABM74126 standard; protein; 34 AA.
XX
AC ABM74126;
XX
DT 17-OCT-2003 (first entry)
DE
DE DNA clone originating in barley containing SNP sequence #536.
XX
KW Barley; single nucleotide polymorphism; SNP; genotype-phenotype analysis.
XX
OS Hordeum vulgare.
XX
PN WO2003057877-A1.
XX

17-JUL-2003.
PD
XX
PF 16-DEC-2002; 2002WO-IB005403.
XX
PR 20-DEC-2001; 2001JP-00387059.
PR 20-DEC-2001; 2001JP-00387131.
PR 20-DEC-2001; 2001JP-00403299.
PR 20-DEC-2001; 2001JP-00403300.
PR 27-SEP-2002; 2002JP-00327515.
XX
FA (UYN-) UNIV JAPAN OKAYAMA.
XX
XX Sato K, Takeda K, Kohara Y;
PI
XX WPI; 2003-587127/55.
XX
XX Single nucleotide polymorphism sites in barley varieties and DNA
PT sequences containing them for analysis and identification of barley
PT varieties and production of barley transformants with desired
PT characteristics.
XX
PS Disclosure; SEQ ID XX; 284pp; Japanese.
XX
XX The present invention relates to oligonucleotide clones originating in
CC barley (Hordeum vulgare) which contain single nucleotide polymorphisms
CC (SNP). The oligonucleotides may be used for analysis of SNPs among barley
CC varieties, identification of particular varieties and genotype-phenotype
CC analysis, isolation of specific genes and creation of new varieties by
CC transformation of barley varieties with them and production of new barley
CC varieties with desired properties. The present sequence represents an
CC oligonucleotide clone sequence featured in the specification. The
CC sequence data for this patent did not form part of the printed
CC specification, but was obtained in electronic format directly from WIPO
CC at ftp.wipo.int/pub/published-pct-sequences
XX
SQ Sequence 34 AA;
Query Match 30.9%; Score 30; DB 7; Length 34;
Best Local Similarity 45.5%; Pred. No. 9.4e+02;
Matches 5; Conservative 3; Mismatches 3; Indels 0; Gaps 0;

QY 7 SKIAFKIVSOE 17
||| |||
Db 7 SKVLFSLVQEE 17

RESULT 93
ADL26414
ID ADL26414 standard; peptide; 37 AA.
XX
AC ADL26414;
XX
DT 17-JUN-2004 (first entry)
DE
DE Synthetic peptide 1553 derived from a conserved region of HCV.
XX
KW HCV; hepatitis C virus; virucide; vaccine; MHC; HLA;
KW major histocompatibility complex; human leukocyte antigen.
XX
OS Synthetic.
XX
PN WO2004024182-A2.
XX
PD 25-MAR-2004.
XX
XX 27-AUG-2003; 2003WO-BP009482.
XX
PR 13-SEP-2002; 2002AT-00001376.
PR 27-FEB-2003; 2003WO-BP002005.
PR 11-JUL-2003; 2003EP-00450171.
XX
PA (INTE-) INTERCELL AG.
XX

```

PI Buschle M, Habel A, Klade C, Mattner F, Otava O, Vytvytska O;  
 XX Zauner W, Zinke S, Kirlappos H;  
 DR WPI; 2004-269899/25.  
 XX Isolating Hepatitis C Virus peptides (Hps) which have a binding capacity  
 PT to a MHC/HLA molecule or a complex comprising the HCV-peptide and the  
 PT molecule by separating the complex from the HCV-peptides which do not  
 XX bind to the molecule.  
 XX Example 1; Page 32; 73pp; English.  
 XX The invention relates to a novel method for isolating Hepatitis C Virus  
 CC (HCV) peptides (Hps). The method of the invention has virucide activity,  
 CC and may be useful in producing a vaccine. The method is useful for  
 CC isolating Hepatitis C Virus peptides (Hps) which have a binding capacity  
 CC to a MHC/HLA molecule or a complex comprising the HCV-peptide and the  
 CC MHC/HLA molecule for preparing a vaccine against HCV infection. The T  
 CC cells, a T cell clone or a T cell population or preparation is useful for  
 CC identifying heteroclitic epitopes or for preparing a composition for  
 CC treating HCV infection. The present sequence represents a synthetic  
 CC peptide derived from a conserved region of HCV.  
 XX Sequence 37 AA;

Query Match 30.9%; Score 30; DB 8; Length 37;  
 Best Local Similarity 66.7%; Pred. No. 1e+03;  
 Matches 6; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

QY 9 IAFKIVSQE 17  
 :|||:|  
 Db 29 VAFKIMSCE 37

## RESULT 94

ADL26416  
 ID ADL26416 standard; peptide; 37 AA.

XX AC ADL26416;

XX DT 17-JUN-2004 (first entry)

XX DE Synthetic peptide 1555 derived from a conserved region of HCV.

XX KW HCV; hepatitis C virus; virucide; vaccine; MHC; HLA;  
 XX major histocompatibility complex; human leukocyte antigen.

XX OS Synthetic.

XX PN WO2004024182-A2.

XX PD 25-MAR-2004.

XX PF 27-AUG-2003; 2003WO-EP009482.

XX PR 13-SEP-2002; 2002AT-00001376.

XX PR 27-FEB-2003; 2003WO-EP002005.

XX PR 11-JUL-2003; 2003EP-00450171.

XX PA (INTE-) INTERCELL AG.

XX PI Buschle M, Habel A, Klade C, Mattner F, Otava O, Vytvytska O;

XX PI Zauner W, Zinke S, Kirlappos H;

XX DR WPI; 2004-269899/25.

XX Isolating Hepatitis C Virus peptides (Hps) which have a binding capacity  
 PT to a MHC/HLA molecule or a complex comprising the HCV-peptide and the  
 PT molecule by separating the complex from the HCV-peptides which do not  
 XX bind to the molecule.

XX Example 1; Page 32; 73pp; English.

XX PS

XX XX

CC The invention relates to a novel method for isolating Hepatitis C Virus  
 CC (HCV) peptides (Hps). The method of the invention has virucide activity,  
 CC and may be useful in producing a vaccine. The method is useful for  
 CC isolating Hepatitis C Virus peptides (Hps) which have a binding capacity  
 CC to a MHC/HLA molecule or a complex comprising the HCV-peptide and the  
 CC MHC/HLA molecule for preparing a vaccine against HCV infection. The T  
 CC cells, a T cell clone or a T cell population or preparation is useful for  
 CC identifying heteroclitic epitopes or for preparing a composition for  
 CC treating HCV infection. The present sequence represents a synthetic  
 CC peptide derived from a conserved region of HCV.  
 XX Sequence 37 AA;

XX Query Match 30.9%; Score 30; DB 8; Length 37;  
 XX Best Local Similarity 66.7%; Pred. No. 1e+03;  
 XX Matches 6; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

QY 9 IAFKIVSQE 17  
 :|||:|  
 Db 29 VAFKIMSCE 37

## RESULT 95

AAW74948  
 ID AAW74948 standard; protein; 39 AA.

XX AC AAW74948;

XX DT 19-JAN-1999 (first entry)

XX DE Human secreted protein encoded by gene 62 clone HATD67.

XX KW Human; secreted protein; testis; tumour; foetal brain tissue;

XX KW fusion protein; cancer; central nervous system; seizure; diagnosis;

XX KW neurodegenerative disease.

XX OS Homo sapiens.

XX PN WO9839448-A2.

XX PD 11-SEP-1998.

XX PF 06-MAR-1998; 98WO-US004493.

XX PR 07-MAR-1997; 97US-0038621P.

XX PR 07-MAR-1997; 97US-0040161P.

XX PR 07-MAR-1997; 97US-0040162P.

XX PR 07-MAR-1997; 97US-0040163P.

XX PR 07-MAR-1997; 97US-0040333P.

XX PR 07-MAR-1997; 97US-0040334P.

XX PR 07-MAR-1997; 97US-0040336P.

XX PR 07-MAR-1997; 97US-0040626P.

XX PR 11-APR-1997; 97US-0043311P.

XX PR 11-APR-1997; 97US-0043312P.

XX PR 11-APR-1997; 97US-0043313P.

XX PR 11-APR-1997; 97US-0043314P.

XX PR 11-APR-1997; 97US-0043315P.

XX PR 11-APR-1997; 97US-0043568P.

XX PR 11-APR-1997; 97US-0043569P.

XX PR 11-APR-1997; 97US-0043576P.

XX PR 11-APR-1997; 97US-0043578P.

XX PR 11-APR-1997; 97US-0043580P.

XX PR 11-APR-1997; 97US-0043669P.

XX PR 11-APR-1997; 97US-0043670P.

XX PR 11-APR-1997; 97US-0043671P.

XX PR 11-APR-1997; 97US-0043672P.

XX PR 11-APR-1997; 97US-0043674P.

XX PR 23-MAY-1997; 97US-0047492P.

XX PR 23-MAY-1997; 97US-0047500P.

XX PR 23-MAY-1997; 97US-0047501P.

XX PR 23-MAY-1997; 97US-0047502P.

XX PR 23-MAY-1997; 97US-0047503P.

XX PR 23-MAY-1997; 97US-0047581P.

PR 23-MAY-1997; 97US-0047582P.  
 PR 23-MAY-1997; 97US-0047583P.  
 PR 23-MAY-1997; 97US-0047584P.  
 PR 23-MAY-1997; 97US-0047585P.  
 PR 23-MAY-1997; 97US-0047586P.  
 PR 23-MAY-1997; 97US-0047587P.  
 PR 23-MAY-1997; 97US-0047588P.  
 PR 23-MAY-1997; 97US-0047589P.  
 PR 23-MAY-1997; 97US-0047590P.  
 PR 23-MAY-1997; 97US-0047592P.  
 PR 23-MAY-1997; 97US-0047593P.  
 PR 23-MAY-1997; 97US-0047594P.  
 PR 23-MAY-1997; 97US-0047595P.  
 PR 23-MAY-1997; 97US-0047596P.  
 PR 23-MAY-1997; 97US-0047597P.  
 PR 23-MAY-1997; 97US-0047598P.  
 PR 23-MAY-1997; 97US-0047599P.  
 PR 23-MAY-1997; 97US-0047600P.  
 PR 23-MAY-1997; 97US-0047601P.  
 PR 23-MAY-1997; 97US-0047612P.  
 PR 23-MAY-1997; 97US-0047613P.  
 PR 23-MAY-1997; 97US-0047614P.  
 PR 23-MAY-1997; 97US-0047615P.  
 PR 23-MAY-1997; 97US-0047616P.  
 PR 23-MAY-1997; 97US-0047617P.  
 PR 23-MAY-1997; 97US-0047618P.  
 PR 23-MAY-1997; 97US-0047632P.  
 PR 23-MAY-1997; 97US-0047633P.  
 PR 06-JUN-1997; 97US-0048964P.  
 PR 06-JUN-1997; 97US-0048974P.  
 PR 13-JUN-1997; 97US-0049610P.  
 PR 08-JUL-1997; 97US-0051526P.  
 PR 16-JUL-1997; 97US-0052874P.  
 PR 18-AUG-1997; 97US-0055724P.  
 PR 22-AUG-1997; 97US-0056630P.  
 PR 22-AUG-1997; 97US-0056631P.  
 PR 22-AUG-1997; 97US-0056632P.  
 PR 22-AUG-1997; 97US-0056633P.  
 PR 22-AUG-1997; 97US-0056637P.  
 PR 22-AUG-1997; 97US-0056662P.  
 PR 22-AUG-1997; 97US-0056664P.  
 PR 22-AUG-1997; 97US-0056845P.  
 PR 22-AUG-1997; 97US-0056862P.  
 PR 22-AUG-1997; 97US-0056864P.  
 PR 22-AUG-1997; 97US-0056872P.  
 PR 22-AUG-1997; 97US-0056874P.  
 PR 22-AUG-1997; 97US-0056875P.  
 PR 22-AUG-1997; 97US-0056876P.  
 PR 22-AUG-1997; 97US-0056877P.  
 PR 22-AUG-1997; 97US-0056878P.  
 PR 22-AUG-1997; 97US-0056879P.  
 PR 22-AUG-1997; 97US-0056880P.  
 PR 22-AUG-1997; 97US-0056881P.  
 PR 22-AUG-1997; 97US-0056882P.  
 PR 22-AUG-1997; 97US-0056883P.  
 PR 22-AUG-1997; 97US-0056884P.  
 PR 22-AUG-1997; 97US-0056886P.  
 PR 22-AUG-1997; 97US-0056887P.  
 PR 22-AUG-1997; 97US-0056888P.  
 PR 22-AUG-1997; 97US-0056889P.  
 PR 22-AUG-1997; 97US-0056892P.  
 PR 22-AUG-1997; 97US-0056893P.  
 PR 22-AUG-1997; 97US-0056894P.  
 PR 22-AUG-1997; 97US-0056903P.  
 PR 22-AUG-1997; 97US-0056908P.  
 PR 22-AUG-1997; 97US-0056909P.  
 PR 22-AUG-1997; 97US-0056910P.  
 PR 22-AUG-1997; 97US-0056911P.  
 PR 05-SEP-1997; 97US-0057650P.  
 PR 05-SEP-1997; 97US-0057669P.  
 PR 12-SEP-1997; 97US-0057761P.  
 PR 12-SEP-1997; 97US-0058785P.  
 PR 02-OCT-1997; 97US-0061060P.  
 PR (HUMA-) HUMAN GENOME SCI INC.

XX Ruben SM, Rosen CA, Fischer CL, Soppet DR, Carter KC,  
 PI Bednarik DP, Endress GA, Yu G, Ni J, Feng P, Young PE, Greene JW,  
 PI Ferrie AM, Duan R, Hu J, Florence KA, Olsen HS, Ebner R, Brewer LA,  
 PI Moore PA, Shi Y, Lafleur DW, Li Y, Zeng Z, Kyaw H;  
 XX WPI; 1998-506364/43.  
 DR N-PSDB; AAV59732.  
 XX  
 DR  
 XX  
 PT New isolated human genes and the secreted polypeptide(s) they encode -  
 PT useful for diagnosis and treatment of e.g. cancers, neurological  
 PT disorders, immune diseases, inflammation or blood disorders.  
 XX  
 PS Claim 1; Page 669; 721pp; English.  
 XX  
 CC This sequence represents a secreted human protein encoded by the nucleic  
 CC acid molecule designated Gene 62 from the human cDNA clone HADPT67  
 CC (deposited as clone ATCC 97900 and ATCC 209046). The gene can be used to  
 CC generate fusion proteins by linking to the gene to a human immunoglobulin  
 CC Fc portion (e.g. AAV59502) for increasing the stability of the fused  
 CC protein as compared to the human protein only. The invention relates to  
 CC 186 novel genes and their fragments (nucleic acid sequences: AAV59511-  
 CC V59812; amino acid sequences AAW74731-W75026) which are useful for  
 CC preventing, treating or ameliorating medical conditions e.g. by protein  
 CC or gene therapy. Also, pathological conditions can be diagnosed by  
 CC determining the amount of the new polypeptides in a sample or by  
 CC determining the presence of mutations in the new polynucleotides.  
 CC Specific uses are described for each of the 186 polynucleotides, based on  
 CC which tissues they are most highly expressed in (see AAV59511 for  
 CC described uses)  
 XX  
 SQ Sequence 39 AA;  
 Query Match 30.9%; Score 30; DB 2; Length 39;  
 Best Local Similarity 58.3%; Pred. No. 1.1e+03;  
 Matches 7; Conservative 1; Mismatches 4; Indels 0; Gaps 0;  
 QY 3 NHLNKKIAPKIV 14  
 DB 27 NHLAPRILFFIV 38  
 RESULT 96  
 AAG74232  
 ID AAG74232 standard; protein; 39 AA.  
 AC AAG74232;  
 XX  
 DT 03-SEP-2001 (first entry)  
 XX  
 DE Human colon cancer antigen protein SEQ ID NO:4996.  
 XX  
 KW Human; colon cancer; colon cancer antigen; diagnosis; detection;  
 KW colorectal carcinoma.  
 XX  
 OS Homo sapiens.  
 XX  
 PN WO200122920-A2.  
 XX  
 PD 05-APR-2001.  
 XX  
 XX 28-SEP-2000; 2000WO-US026524.  
 PF  
 XX 29-SEP-1999; 99US-0157137P.  
 PR 03-NOV-1999; 99US-0163280P.  
 PR  
 XX (HUMA-) HUMAN GENOME SCI INC.  
 PA  
 XX Ruben SM, Barash SC, Birse CE, Rosen CA;  
 PI WPI; 2001-235357/24.  
 DR N-PSDB; AAV33663.  
 DR  
 XX

Nucleic acids encoding 4277 human colon cancer-associated polypeptides, useful for preventing, diagnosing and/or treating colorectal cancers.

Claim 11; Page 6739-6740; 9803pp; English.

AAH37195 and AAG73514 to AAG77788 represent human colon cancer-associated nucleic acid molecules (N) and proteins (P), where the proteins are collectively known as colon cancer antigens. The colon cancer antigens have cytostatic activity and can be used in gene therapy and vaccine production. N and P may be used in the prevention, diagnosis and treatment of diseases associated with inappropriate P expression. For example, N and P may be used to treat disorders associated with decreased expression by rectifying mutations or deletions in a patient's genome that affect the activity of P by expressing inactive proteins or to supplement the patient's own production of P. Additionally, N may be used to produce the colon cancer-associated Ps, by inserting the nucleic acids into a host cell and culturing the cell to express the proteins. N and P can be used in the prevention, diagnosis and treatment of colorectal carcinomas and cancers. AAH37196 to AAH37204 and AAG77789 represent sequences used in the exemplification of the present invention. N.B. Pages 666 to 682 and page 7053 of the sequence listing were missing at time of publication, meaning no sequences are present for SEQ ID NO:1027 to 1052, 7921 and 7922

```

AA      SQ      Sequence 39 AA;
Query Match      30.3%; Score 30; DB 4; Length 39;
Best Local Similarity 42.9%; Pred.No. 1.1e+03;
Matches 6: Conservative 4; Mismatches 4; Indels 0; Gaps 0;

```

Qy 1 EPNHLSKIAFKIV 14  
:|::|||:|  
Dp 19 KPHYLNIKLPPNIV 32

RESULT 97  
ABG95405  
ID ABG95405 standard: protein: 39 AA:

AC ABG95405:

DT 15-JAN-2003 (first entry)

Human novel secreted protein #226.

Human; secreted protein; autoimmune disease; chemotaxis; rheumatoid arthritis; hyperproliferative disorder; breast neoplasm; liver neoplasm cardiovascular disorder; cardiac arrest; skin aging; cerebrovascular disorder; cerebral ischaemia; angiogenesis; sunburn; nervous system disorders; Alzheimer's disease; infection; corneal disorder; corneal infection; wound healing; tissue regeneration; epithelial cell proliferation; organ transplantation; food additive; preservative; nutritional

AA  
OS Homo sapiens.

XX  
PN  
US6420526-B1.

16-JUL-2002.

08-SEP-1998;

07-MAR-1997;

PR 07-MAR-1997;

PR 07-MAR-1997;

PR 07-MAR-1997;

PR 11-APR-1997;

PR 11-APR-1997;

[illegible]

XX	18-JUN-2002	(first entry)	Insulin/insulin-like growth factor receptor-binding peptide #686.
XX			Cytostatic; antidiabetic; neuroprotective; cerebroprotective; ophthalmological; insulin; receptor; gene therapy; diabetes; insulin-like growth factor-1; IGF-1; tumour; prostate; breast; diabetic retinopathy; neurological diseases; stroke; diabetic neuropathy.
XX			Synthetic.
XX	W0200172771-A2.		
XX	04-OCT-2001.		
XX	29-MAR-2000;	2000WO-US008528.	
XX	29-MAR-2000;	2000WO-US008528.	
XX	(DGIB-) DGI BIOTECHNOLOGIES LLC.		
XX	(NOVO) NOVO NORDISK AS.		
XX	Beasley J, Blume AJ, Schaeffer L, Pillutia R, Brandt J; Brissette R, Spetzler J, Cheng W, Ostergaard S, Manddecki WS; Hansen PH, Ravera M, Hsiao K;		
XX	WPI; 2002-025774/03.		
XX	Modulating insulin activity in mammalian cells, for treating e.g. diabetes and tumors, comprises using peptides that bind to insulin or insulin-like growth factor receptors.		
XX	Disclosure; Fig 1G-2; 390pp; English.		
XX	The invention relates to a method of modulating insulin activity in mammalian cells by administering a peptide that binds the insulin receptor (IR). A composition containing a peptide, optionally expressed from gene therapy vectors, that binds to Site 1 of IR and an insulin agonist are useful for treating diabetes. Also, peptides that are antagonists of the insulin-like growth factor-1 (IGF-1) receptor are useful for treating insulin-like growth factor-1 (IGF-1)-sensitive tumours (e.g. of prostate and breast) and diabetic retinopathy, while IGF-1 receptor agonists are useful for treating neurological diseases, including stroke and diabetic neuropathy. The peptides are also useful in screening for compounds that bind to IR or IGF-1 receptor, potential therapeutics and research reagents. AAU8034-AAU90957 represent IR and/or IGF-1 receptor-binding peptides and related amino acid sequences of the invention		
XX	Sequence 39 AA;		
XX	Query Match	30.9%;	Score 30; DB 5; Length 39;
XX	Best Local Similarity	60.0%;	Pred. No. 1.1e+03;
XX	Matches	6; Conservative	2; Mismatches 2; Indels 0; Gaps 0;
XX	QY	10 AFKIVSQEPPA 19	
XX	Db	14 ASKVSEPPA 23	
XX	RESULT 99		
XX	ADA03571		
XX	ID	ADA03571 standard; peptide; 39 AA.	
XX	AC	ADA03571;	
XX	XX		
XX	DT	06-NOV-2003	(first entry)
XX			Insulin receptor (IR) related Formula 1 motif peptide SEQ ID NO:202.
XX	DE		insulin-like growth factor receptor; IGFR; modulate; insulin receptor;
XX	KW		IR; insulin; cycostatic; IGFR agonist; IGFR antagonist; cancer;

KW Leukaemia; sarcoma; lymphoma; carcinoma.  
 XX  
 OS Synthetic.  
 OS Homo sapiens.  
 XX  
 PN WC2003027246-A2.  
 XX  
 XX  
 XX  
 PD 03-APR-2003.  
 XX  
 PF 24-SEP-2002; 2002WO-US030412.  
 XX  
 XX 24-SEP-2001; 2001US-00962756.  
 XX  
 XX (NOVO ) NOVO NORDISK AS.  
 PA (DGIB-) DGI BIOTECHNOLOGIES.  
 XX  
 XX Pillutla R, Dedova O, Blume AJ, Goldstein NI, Brissette R;  
 PI Wang P, Liu H, Hsiao K, Lennick M, Fletcher P;  
 PI WPI; 2003-363211/34.  
 XX  
 XX Modulating insulin-like growth factor receptor (IGFR) activity in IGF-  
 PT responsive mammalian cells, useful for treating cancer comprises  
 PT contacting the cells with an amino acid sequence to modulate the activity  
 PT of IGFR.  
 XX  
 XX Disclosure; Fig 1G; 372pp; English.  
 XX  
 CC The present invention describes a method for modulating insulin-like  
 CC growth factor receptor (IGFR) activity in insulin-like growth factor-  
 CC responsive mammalian cells comprising contacting the cells with an amino  
 CC acid sequence to modulate the activity of IGFR. In modulating IGFR  
 CC activity, the amino acid sequence comprises XIX2X3X4X5 (1), where X1, X2  
 CC and X5 = phenylalanine or tyrosine; X3 = aspartic acid, glutamic acid,  
 CC glycine or serine; and X4 = tryptophan, tyrosine or phenylalanine. The  
 CC amino acid sequence is not insulin, insulin-like growth factor, an anti-  
 CC insulin receptor antibody, an anti-insulin-like growth receptor antibody,  
 CC or its fragment. Also described: (1) decreasing or increasing IGFR  
 CC activity in IGF-responsive mammalian cells by contacting the cells with  
 CC an amino acid sequence to decrease or increase the activity of IGFR; (2)  
 CC an IGF modulator, agonist or antagonist; (3) identifying IGF modulator;  
 CC and (4) enhancing survival of an IGF-responsive mammalian cell by  
 CC contacting the cell with (1) to enhance the survival of the cell. IGFR  
 CC modulators have cytostatic activity, and can be used as IGF agonists or  
 CC IGF antagonists. The methods, modulators, agonists and antagonists are  
 CC useful for treating cancer, e.g. leukaemia, sarcoma, lymphoma or  
 CC carcinoma. The methods are useful for identifying molecular structures  
 CC that are capable of acting as an IGF agonist or antagonist. The present  
 CC sequence represents a peptide given in the exemplification of the present  
 CC invention.  
 XX  
 XX Sequence 39 AA;  
 SQ  
 Query Match 30.9%; Score 30; DB 6; Length 39;  
 Best Local Similarity 60.0%; Pred. No. 1.1e+03;  
 Matches 6; Conservative 2; Mismatches 2; Indels 0; Gaps 0;  
 QY 10 AFKIVSQEPA 19  
 DB 14 ASKVSEPPA 23  
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 RESULT 100  
 ABO34599  
 ID ABO34599 standard; protein; 39 AA.  
 XX  
 AC ABO34599;  
 XX  
 XX 22-SEP-2003 (first entry)  
 DT  
 XX Region of human secreted protein encoded by cDNA sequence #226.  
 DE  
 XX Human; secreted protein; hyperproliferative disorder; leukaemia;  
 KW

KW breast cancer; wound; reproductive disorder; blood-related disorder;  
 KW haemophilia; thrombocytopaenia; immunodeficiency; thymic hypoplasia;  
 KW Wiskott-Aldrich syndrome; autoimmune disorder; multiple sclerosis;  
 KW graft-versus-host disease; Hashimoto's thyroiditis; allergy; asthma;  
 KW viral infection; bacterial infection; fungal infection; AIDS; sepsis;  
 KW renal disorder; kidney failure; cardiovascular disorder; cytostatic;  
 KW angina pectoris; cerebral ischaemia; congenital heart defect;  
 KW respiratory disorder; neurological disorder; Alzheimer's disease;  
 KW Parkinson's disease; inflammation; Crohn's disease; vulvular;  
 KW immunosuppressive; antibacterial; haemostatic; thrombolytic;  
 KW anticoagulant; neuroprotective; thyromimetic; antiallergic;  
 KW antiasthmatic; virucide; fungicide; anti-HIV; nephrotropic; antiangiinal;  
 KW cerebroprotective; cardiac; nootropic; antiparkinsonian;  
 XX  
 XX antiinflammatory.  
 OS Homo sapiens.  
 XX  
 PN US2003049618-A1.  
 XX  
 XX 13-MAR-2003.  
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 XX  
 XX 16-MAR-2001; 2001US-00809391.  
 PF  
 XX 07-MAR-1997; 97US-0038621P.  
 XX 07-MAR-1997; 97US-0040162P.  
 PR 07-MAR-1997; 97US-0040163P.  
 PR 07-MAR-1997; 97US-0040333P.  
 PR 07-MAR-1997; 97US-0040334P.  
 PR 07-MAR-1997; 97US-0040336P.  
 PR 07-MAR-1997; 97US-0040626P.  
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 PR 23-MAY-1997; 97US-0047612P.  
 PR 23-MAY-1997; 97US-0047613P.  
 PR 23-MAY-1997; 97US-0047614P.



PR 23-MAY-1997; 97US-0047615P.  
 PR 23-MAY-1997; 97US-0047617P.  
 PR 23-MAY-1997; 97US-0047618P.  
 PR 23-MAY-1997; 97US-0047632P.  
 PR 23-MAY-1997; 97US-0047633P.  
 PR 06-JUN-1997; 97US-0048964P.  
 PR 13-JUN-1997; 97US-0048974P.  
 PR 08-JUL-1997; 97US-0049610P.  
 PR 16-JUL-1997; 97US-0051926P.  
 PR 18-AUG-1997; 97US-0052874P.  
 PR 22-AUG-1997; 97US-0055724P.  
 PR 22-AUG-1997; 97US-0056630P.  
 PR 22-AUG-1997; 97US-0056631P.  
 PR 22-AUG-1997; 97US-0056632P.  
 PR 22-AUG-1997; 97US-0056633P.  
 PR 22-AUG-1997; 97US-0056637P.  
 PR 22-AUG-1997; 97US-0056662P.  
 PR 22-AUG-1997; 97US-0056664P.  
 PR 22-AUG-1997; 97US-0056845P.  
 PR 22-AUG-1997; 97US-0056852P.  
 PR 22-AUG-1997; 97US-0056862P.  
 PR 22-AUG-1997; 97US-0056864P.  
 PR 22-AUG-1997; 97US-0056872P.  
 PR 22-AUG-1997; 97US-0056874P.  
 PR 22-AUG-1997; 97US-0056875P.  
 PR 22-AUG-1997; 97US-0056876P.  
 PR 22-AUG-1997; 97US-0056877P.  
 PR 22-AUG-1997; 97US-0056878P.  
 PR 22-AUG-1997; 97US-0056879P.  
 PR 22-AUG-1997; 97US-0056880P.  
 PR 22-AUG-1997; 97US-0056881P.  
 PR 22-AUG-1997; 97US-0056882P.  
 PR 22-AUG-1997; 97US-0056884P.  
 PR 22-AUG-1997; 97US-0056886P.  
 PR 22-AUG-1997; 97US-0056887P.  
 PR 22-AUG-1997; 97US-0056888P.  
 PR 22-AUG-1997; 97US-0056889P.  
 PR 22-AUG-1997; 97US-0056892P.  
 PR 22-AUG-1997; 97US-0056893P.  
 PR 22-AUG-1997; 97US-0056894P.  
 PR 22-AUG-1997; 97US-0056903P.  
 PR 22-AUG-1997; 97US-0056908P.  
 PR 22-AUG-1997; 97US-0056909P.  
 PR 22-AUG-1997; 97US-0056910P.  
 PR 22-AUG-1997; 97US-0056911P.  
 PR 05-SEP-1997; 97US-0057650P.  
 PR 05-SEP-1997; 97US-0057669P.  
 PR 12-SEP-1997; 97US-0057761P.  
 PR 09-OCT-1997; 97US-0058785P.  
 PR 06-MAR-1998; 97US-0061660P.  
 PR 08-SEP-1998; 98MO-US004493.  
 PR 17-MAR-2000; 98US-0019476.  
 PR 2000US-0190068P.

XX (RUBE/) RUBEN S M.  
 PA (ROSE/) ROSEN C A.  
 PA (SOPP/) SOPPET D R.  
 PA (CART/) CARTER K C.  
 PA (BEDN/) BEDNARIK D P.  
 PA (ENDR/) ENDRESS G A.  
 PA (YUG/) YU G.  
 PA (NIJ/) NI J.  
 PA (FENG/) FENG P.  
 PA (YOUN/) YOUNG P E.  
 PA (GREE/) GREENE J M.  
 PA (FERR/) FERRIE A M.  
 PA (DUAN/) DUAN D R.  
 PA (HUJ/) HU J.  
 PA (FLOR/) FLORENCE K A.  
 PA (OLSE/) OLSEN H S.  
 PA (FISC/) FISCHER C L.  
 PA (EBNE/) EBNER R L.  
 PA (BREW/) BREWER L A.  
 PA (MOOR/) MOORE P A.

PA (SHLY/) SHI Y.  
 PA (LAPL/) LAFLEUR D W.  
 PA (LIYY/) LI Y.  
 PA (ZENG/) ZENG Z.  
 PA (KYAW/) KYAW H.

PI Ruben SM, Rosen CA, Soppet DR, Carter KC, Bednariak DP;  
 PI Endress GA, Yu G, Ni J, Feng P, Greene JM, Ferrie AM;  
 PI Duan DR, Hu J, Florence KA, Olsen HS, Fischer CL, Ebner R;  
 PI Brewer LA, Moore PA, Shi Y, Lafleur DW, Li Y, Zeng Z, Kyaw H;  
 XX WPI: 2003-521800/49.  
 DR N-PSDB: ACD82866.

XX New genes and its encoded prostate cancer antigen proteins, useful for  
 PT preventing, treating, ameliorating or diagnosing e.g. prostate cancers,  
 PT thymic hypoplasia, multiple sclerosis, AIDS, angina pectoris or cerebral  
 PT ischemia.

PS Claim 3; SEQ ID NO 545; 260pp; English.

XX The present invention relates to the isolation of novel human secreted  
 CC proteins and the polynucleotide sequences encoding them. The invention  
 CC also discloses vectors, host cells, antibodies, and recombinant methods  
 CC for producing human secreted proteins. The polypeptide and polynucleotide  
 CC sequences for the secreted proteins are useful for preventing, treating,  
 CC ameliorating or diagnosing medical conditions such as hyperproliferative  
 CC disorders (e.g. leukaemia or breast cancers), wounds, reproductive  
 CC disorders, blood-related disorders (e.g. haemophilia or  
 CC thrombocytopaenia), immunodeficiencies (e.g. Wiskott-Aldrich syndrome or  
 CC thymic hypoplasia), autoimmune disorders (e.g. graft-versus-host disease,  
 CC multiple sclerosis or Hashimoto's thyroiditis), allergies (e.g. asthma),  
 CC viral or bacterial or fungal infections (e.g. AIDS or sepsis), renal  
 CC disorders (e.g. kidney failure), cardiovascular disorders (e.g. angina  
 CC pectoris, cerebral ischaemia or congenital heart defects), respiratory  
 CC disorders, neurological disorders (e.g. Alzheimer's disease or  
 CC Parkinson's disease), and inflammations (e.g. Crohn's disease). The  
 CC polynucleotide or polypeptide may also be used as vaccine adjuvants.  
 CC ABO34374-ABO34815 represent human secreted proteins or their fragments.  
 CC Note: The sequence data for this patent did not form part of the printed  
 CC specification, but was obtained in electronic format directly from the  
 CC USPTO web site at seqdata.uspto.gov/psipdIDEntry.html

SQ Sequence 39 AA;

Query Match 30.9%; Score 30; DB 6; Length 39;  
 Best Local Similarity 58.3%; Pred. No. 1.1e+03;  
 Matches 7; Conservative 1; Mismatches 4; Indels 0; Gaps 0;

QY 3 NHLSKIAFKIV 14  
 Db ||||:||||  
 27 NHLAFRILFFIV 38

RESULT 101  
 ADH94784  
 ID ADH94784 standard; protein; 39 AA.

XX AC ADH94784;

XX 22-APR-2004 (first entry)

XX Insulin receptor motif polypeptide, SEQ ID No 202.

XX insulin receptor; insulin; insulin-like growth factor receptor; agonist;  
 KW antagonist; antidiabetic; diabetes; insulin shock.

XX Unidentified.

XX WO2003070747-A2.

XX 28-AUG-2003.

PF 24-SEP-2002; 2002WO-US030312.  
XX  
PR 24-SEP-2001; 2001US-00962756.  
XX  
XX (NOVO ) NOVO NORDISK AS.  
PA (DGIB-) DGI BIOTECHNOLOGIES.  
XX  
XX Pillutla R, Brissette R, Blume AJ, Schaeffer L, Brandt J;  
PI Goldstein NI, Spetzler J, Ostergaard S;  
XX MPI; 2003-833235/77.  
DR  
XX  
XX Modulating insulin-like growth factor receptor (IGFR) activity in IGF-  
PT responsive mammalian cells, useful for treating diabetes comprises  
PT contacting the cells with an amino acid sequence to modulate the activity  
PT of IGFR.  
XX  
XX Claim 7; SEQ ID NO 202; 328pp; English.  
PS  
XX The invention relates to a novel method for decreasing or increasing  
CC insulin receptor activity in mammalian cells. The invention further  
CC relates to peptide sequences capable of binding to insulin and/or insulin  
CC -like growth factor receptors with either agonist or antagonist activity. The  
CC peptide sequences are identified from various peptide libraries. The  
CC novel method comprises administering to the mammalian cells an amino acid  
CC having subsequences that binds to site 1 and site 2 of an insulin  
CC receptor. The subsequences are joined C-terminus to N-terminus and  
CC oriented site 1 to site 2. The sequence is not insulin or insulin-like  
CC growth factor. The peptide sequences of the invention have antidiabetic  
CC activity. The peptides are useful for treating diabetes or insulin shock.  
CC This sequence represents an insulin receptor/ insulin growth factor  
CC receptor binding polypeptide relating to the invention.  
XX  
XX Sequence 39 AA;  
SQ  
Query Match 30.9%; Score 30; DB 7; Length 39;  
Best Local Similarity 60.0%; Pred No. 1.1e+03;  
Matches 6; Conservative 2; Mismatches 2; Indels 0; Gaps 0;  
Qy 10 AFKIVSQEPA 19  
Db 14 ASKVSEPPA 23  
RESULT 102  
AD123260  
ID AD123260 standard; protein; 39 AA.  
XX  
AC AD123260;  
XX  
DT 22-APR-2004 (first entry)  
XX  
XX Novel human secreted protein seq id 545.  
DE  
XX cytotostatic; gene therapy; cancer; human; secreted protein.  
KW  
XX Homo sapiens.  
OS  
XX US2003175858-A1.  
PN  
XX 18-SEP-2003.  
PD  
XX 18-JUN-2001; 2001US-00882171.  
PF  
XX 07-MAR-1997; 97US-0038621P.  
PR 07-MAR-1997; 97US-0040162P.  
PR 07-MAR-1997; 97US-0040163P.  
PR 07-MAR-1997; 97US-0040333P.  
PR 07-MAR-1997; 97US-0040334P.  
PR 07-MAR-1997; 97US-0040336P.  
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PR 11-APR-1997; 97US-0043670P.  
PR 11-APR-1997; 97US-0043671P.  
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PR 11-APR-1997; 97US-0043674P.  
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PR 06-JUN-1997; 97US-0048964P.  
PR 06-JUN-1997; 97US-0048974P.  
PR 13-JUN-1997; 97US-0049610P.  
PR 08-JUL-1997; 97US-0051926P.  
PR 16-JUL-1997; 97US-0052874P.  
PR 18-AUG-1997; 97US-0055724P.  
PR 22-AUG-1997; 97US-0056630P.  
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PR 22-AUG-1997; 97US-0056874P.  
PR 22-AUG-1997; 97US-0056875P.  
PR 22-AUG-1997; 97US-0056876P.  
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PR 22-AUG-1997; 97US-0056886P.  
 PR 22-AUG-1997; 97US-0056887P.  
 PR 22-AUG-1997; 97US-0056888P.  
 PR 22-AUG-1997; 97US-0056889P.  
 PR 22-AUG-1997; 97US-0056892P.  
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 PR 22-AUG-1997; 97US-0056910P.  
 PR 05-SEP-1997; 97US-0057650P.  
 PR 05-SEP-1997; 97US-0057669P.  
 PR 12-SEP-1997; 97US-0057761P.  
 PR 09-OCT-1997; 97US-0058785P.  
 PR 06-MAR-1998; 97US-0061660P.  
 PR 08-SEP-1998; 98WO-US004493.  
 PR 17-MAR-2000; 2000US-0190068P.  
 PR 16-MAR-2001; 2001US-00809391.  
 XX

(RUBE/) RUBEN S M.  
 PA (ROSE/) ROSEN C A.  
 PA (SOPP/) SOPPET D R.  
 PA (CART/) CARTER K C.  
 PA (BEDN/) BEDNARIK D P.  
 PA (ENDR/) ENDRESS G A.  
 PA (YUGG/) YU G.  
 PA (NIJJ/) NI J.  
 PA (FENG/) FENG P.  
 PA (YOUN/) YOUNG P E.  
 PA (GREE/) GREENE J M.  
 PA (FERR/) FERRIE A M.  
 PA (DUAN/) DUAN D R.  
 PA (HULJ/) HU J.  
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 PA (OLSE/) OLSEN H S.  
 PA (FISC/) FISCHER C L.  
 PA (EBNE/) EBNER R.  
 PA (BREW/) BREWER L A.  
 PA (MOOR/) MOORE P A.  
 PA (SHIY/) SHI Y.  
 PA (LAFLE/) LAFLEUR D W.  
 PA (LIYY/) LI Y.  
 PA (ZENG/) ZENG Z.  
 PA (KYAW/) KYAW H.  
 XX

Ruben SM, Rosen CA, Soppet DR, Carter KC, Bednarik DP;  
 Endress GA, Yu G, Ni J, Feng P, Young PE, Greene JM, Ferrie AM;  
 Duan DR, Hu J, Florence KA, Olsen HS, Fischer CL, Ebner R;  
 Brewer LA, Moore PA, Shi Y, Lafleur DW, Li Y, Zeng Z, Kyaw H;

WPI: 2003-898535/82.  
 N-PSDB; ADI22951.

New nucleic acid molecule, useful for preparing a medicament for  
 diagnosing, preventing, treating or ameliorating a medical condition  
 e.g., cancer.

Claim 11; SEQ ID NO 545; 256pp; English.

The invention describes an isolated nucleic acid comprising a sequence  
 having 95 % identity with: a polynucleotide fragment of a sequence not  
 given in the specification, or its allelic variant; a polynucleotide  
 fragment of the cDNA sequence; a polynucleotide sequence encoding a  
 polypeptide, or its fragment, domain, epitope or species homologue; or a  
 polynucleotide that hybridises under stringent conditions to any one of  
 the sequences of (a)-(c). The nucleic acid is useful for preparing a  
 medicament for diagnosing, preventing, treating or ameliorating a medical  
 condition e.g., cancer. The is the amino acid sequence of a novel human  
 secreted protein of the invention.

SQ Sequence 39 AA;

Query March 30.9%; Score 30; DB 7; Length 39;  
 Best Local Similarity 58.3%; Pred. No. 1.1e+03;  
 Matches 7; Conservative 1; Mismatches 4; Indels 0; Gaps 0;

Qy 3 NHLNSKIAFKIV 14  
 |||||:|:|:  
 Db 27 NHLAFRLFFIV 38

RESULT 103

ADH74262

ID ADH74262 standard; protein; 39 AA.

XX ADH74262;

XX DT 25-MAR-2004 (first entry)

XX XX Human secreted protein #226.

XX human; secreted protein; cancer; haematopoietic disorder;  
 KW endocrine disorder; immune system disease; inflammatory disorder.  
 XX Homo sapiens.

XX US2003225248-A1.

XX PD 04-DEC-2003.

XX 10-JUN-2002; 2002US-00164861.

XX 07-MAR-1997; 97US-0038621P.

PR 07-MAR-1997; 97US-0040161P.

PR 07-MAR-1997; 97US-0040162P.

PR 07-MAR-1997; 97US-0040163P.

PR 07-MAR-1997; 97US-0040333P.

PR 07-MAR-1997; 97US-0040334P.

PR 07-MAR-1997; 97US-0040336P.

PR 11-APR-1997; 97US-0043311P.

PR 11-APR-1997; 97US-0043312P.

PR 11-APR-1997; 97US-0043313P.

PR 11-APR-1997; 97US-0043314P.

PR 11-APR-1997; 97US-0043315P.

PR 11-APR-1997; 97US-0043568P.

PR 11-APR-1997; 97US-0043569P.

PR 11-APR-1997; 97US-0043576P.

PR 11-APR-1997; 97US-0043578P.

PR 11-APR-1997; 97US-0043580P.

PR 11-APR-1997; 97US-0043669P.

PR 11-APR-1997; 97US-0043670P.

PR 11-APR-1997; 97US-0043671P.

PR 11-APR-1997; 97US-0043672P.

PR 11-APR-1997; 97US-0043674P.

PR 23-MAY-1997; 97US-0047492P.

PR 23-MAY-1997; 97US-0047500P.

PR 23-MAY-1997; 97US-0047501P.

PR 23-MAY-1997; 97US-0047502P.

PR 23-MAY-1997; 97US-0047503P.

PR 23-MAY-1997; 97US-0047581P.

PR 23-MAY-1997; 97US-0047582P.

PR 23-MAY-1997; 97US-0047583P.

PR 23-MAY-1997; 97US-0047584P.

PR 23-MAY-1997; 97US-0047585P.

PR 23-MAY-1997; 97US-0047586P.

PR 23-MAY-1997; 97US-0047587P.

PR 23-MAY-1997; 97US-0047588P.

PR 23-MAY-1997; 97US-0047589P.

PR 23-MAY-1997; 97US-0047590P.

PR 23-MAY-1997; 97US-0047592P.

PR 23-MAY-1997; 97US-0047593P.

PR 23-MAY-1997; 97US-0047594P.



PA (LIUH/) LIU H.  
 PA (HSIA/) HSIAO K.  
 PA (LENN/) LENNICK M.  
 PA (FLET/) FLETCHER P.  
 XX  
 XX  
 PI Pillutla R, Dedova O, Blume AJ, Goldstein NI, Brissette R;  
 PI Wang P, Liu H, Hsiao K, Lennick M, Fletcher P;  
 XX  
 XX  
 DR WPI: 2004-132606/13.  
 XX  
 PS Disclosure; SEQ ID NO 202; 242pp; English.  
 XX  
 CC The invention describes the use of molecular structures, preferably  
 CC peptides for modulating, increasing or decreasing insulin-like growth  
 CC factor receptor activity in insulin-like growth factor-responsive  
 CC mammalian cells. Also described are: modulating, decreasing or increasing  
 CC insulin-like growth factor receptor activity in insulin-like growth  
 CC factor-responsive mammalian cells; an insulin-like growth factor receptor  
 CC modulator; an insulin-like growth factor receptor antagonist; an insulin-  
 CC like growth factor receptor agonist; identifying an insulin-like growth  
 CC factor receptor modulator; and enhancing survival of an insulin-like  
 CC growth factor-responsive mammalian cell. The molecular structures of the  
 CC peptides are useful for modulating, increasing or decreasing insulin-like  
 CC growth factor receptor activity in insulin-like growth factor-responsive  
 CC mammalian cells. The peptide sequences are useful for binding to insulin  
 CC and/or insulin-like growth factor receptors with either agonist or  
 CC antagonist activity. As agonists, the peptides are useful for development  
 CC of therapeutics to supplement or replace endogenous peptide hormones. The  
 CC antagonist peptides can also be developed as therapeutics. The IR and IGF  
 CC -1R agonist and antagonist peptides are useful as lead compounds for  
 CC identifying other more potent or selective therapeutics, assay reagents  
 CC for identifying other useful ligands, as research tools for further  
 CC analysis of IR and IGF-1R. The IGF-1R antagonists are useful as treatment  
 CC for cancer, e.g. breast, prostate, colorectal or ovarian cancer. This is  
 CC the amino acid sequence of a IGF-1R binding peptide obtained by  
 CC panning a peptide library against the insulin-like growth factor 1  
 CC receptor (IGF-1R) and insulin receptor (IR). Note: This sequence is also  
 CC available in electronic format from the US patent office at  
 CC ftp.segdata.uspto.gov/sequence.html?DocID=20040023887.  
 XX  
 XX Sequence 39 AA;

Query Match 30.9%; Score 30; DB 8; Length 39;  
 Best Local Similarity 60.0%; Pred. No. 1.1e+03;  
 Matches 6; Conservative 2; Mismatches 2; Indels 0; Gaps 0;

OY 10 AFKIVSOEPA 19  
 DB 14 ASKVSEPPA 23

RESULT 105  
 ADM37320  
 ID ADM37320 standard; peptide; 39 AA.  
 XX  
 AC ADM37320;  
 XX

DT 03-JUN-2004 (first entry)  
 DE Anti-IR formula 1 motif peptide #125.  
 XX  
 KW insulin receptor activity; insulin-related disease;  
 KW insulin-like growth factor-related disease; diabetes; insulin shock.  
 XX  
 OS Synthetic.  
 XX

PN US2003236190-A1.  
 XX  
 XX 25-DEC-2003.  
 XX

PF 24-SEP-2002; 2002US-00253471.  
 XX  
 XX 02-SEP-1998; 98US-00146127.  
 PR

PR 29-MAR-2000; 2000US-00538038.  
 PR 24-SEP-2001; 2001US-00962756.

PA (PILL/) PILLUTLA R.  
 PA (BRIS/) BRISSETTE R.  
 PA (BLUM/) BLUME A J.  
 PA (SCHA/) SCHAEFFER L.  
 PA (BRAN/) BRANDT J.  
 PA (GOLD/) GOLDSTEIN N I.  
 PA (SPET/) SPETZLER J.  
 PA (OSTE/) OSTERGAARD S.  
 PA (HAN/) HANSEN P H.

PI Pillutla R, Brissette R, Blume AJ, Schaffer L, Brandt J;  
 PI Goldstein NI, Spetzler J, Ostergaard S, Hansen PH;  
 XX  
 XX WPI: 2004-081583/08.

XX Decreasing or increasing insulin receptor activity in mammalian cells  
 XX using peptide sequences that bind insulin and/or insulin-like growth  
 XX factor receptors, useful for treating diabetes and insulin shock.

XX Claim 7; SEQ ID NO 202; 203pp; English.

XX The invention relates to a method of decreasing insulin receptor activity  
 XX in mammalian cells which comprises administering an amino acid sequence  
 XX having a subsequence comprising a sequence that binds to Site 1 and Site  
 XX 2 of insulin receptor, where the subsequences are linked C-terminus to N-  
 XX terminus and oriented Site 1 to Site 2, where the amino acid sequence is  
 XX not insulin, insulin-like growth factor or their fragments. The methods  
 XX and compositions of the present invention are useful for treating insulin  
 XX - or insulin-like growth factor-related diseases or conditions, such as  
 XX diabetes and insulin shock. The present sequence is used in the  
 XX exemplification of the present invention.

XX Sequence 39 AA;

Query Match 30.9%; Score 30; DB 8; Length 39;  
 Best Local Similarity 60.0%; Pred. No. 1.1e+03;  
 Matches 6; Conservative 2; Mismatches 2; Indels 0; Gaps 0;

OY 10 AFKIVSOEPA 19  
 DB 14 ASKVSEPPA 23

RESULT 106  
 AAW74791  
 ID AAW74791 standard; protein; 40 AA.  
 XX  
 AC AAW74791;

DT 25-JAN-1999 (first entry)  
 XX  
 DE Human secreted protein encoded by gene 62 clone HATDT67.

XX Human; secreted protein; testis; tumour; foetal brain tissue;  
 XX fusion protein; cancer; central nervous system; seizure; diagnosis;  
 XX neurodegenerative disease.

OS Homo sapiens.

XX Key Location/Qualifiers  
 XX FT Misc-difference 40  
 XX /label= unknown

PN WO9839448-A2.

XX 11-SEP-1998.

XX 06-MAR-1998; 98WO-US004493.

XX 07-MAR-1997; 97US-0038621P.

```

PR 07-MAR-1997; 97US-0040161P.
PR 07-MAR-1997; 97US-0040162P.
PR 07-MAR-1997; 97US-0040163P.
PR 07-MAR-1997; 97US-0040333P.
PR 07-MAR-1997; 97US-0040334P.
PR 07-MAR-1997; 97US-0040336P.
PR 07-MAR-1997; 97US-0040326P.
PR 11-APR-1997; 97US-0043311P.
PR 11-APR-1997; 97US-0043312P.
PR 11-APR-1997; 97US-0043313P.
PR 11-APR-1997; 97US-0043314P.
PR 11-APR-1997; 97US-0043315P.
PR 11-APR-1997; 97US-0043568P.
PR 11-APR-1997; 97US-0043569P.
PR 11-APR-1997; 97US-0043576P.
PR 11-APR-1997; 97US-0043580P.
PR 11-APR-1997; 97US-0043578P.
PR 11-APR-1997; 97US-0043659P.
PR 11-APR-1997; 97US-0043670P.
PR 11-APR-1997; 97US-0043671P.
PR 11-APR-1997; 97US-0043672P.
PR 11-APR-1997; 97US-0043674P.
PR 23-MAY-1997; 97US-0047492P.
PR 23-MAY-1997; 97US-0047500P.
PR 23-MAY-1997; 97US-0047501P.
PR 23-MAY-1997; 97US-0047502P.
PR 23-MAY-1997; 97US-0047503P.
PR 23-MAY-1997; 97US-0047581P.
PR 23-MAY-1997; 97US-0047582P.
PR 23-MAY-1997; 97US-0047583P.
PR 23-MAY-1997; 97US-0047584P.
PR 23-MAY-1997; 97US-0047585P.
PR 23-MAY-1997; 97US-0047586P.
PR 23-MAY-1997; 97US-0047587P.
PR 23-MAY-1997; 97US-0047588P.
PR 23-MAY-1997; 97US-0047589P.
PR 23-MAY-1997; 97US-0047590P.
PR 23-MAY-1997; 97US-0047592P.
PR 23-MAY-1997; 97US-0047593P.
PR 23-MAY-1997; 97US-0047594P.
PR 23-MAY-1997; 97US-0047595P.
PR 23-MAY-1997; 97US-0047596P.
PR 23-MAY-1997; 97US-0047597P.
PR 23-MAY-1997; 97US-0047598P.
PR 23-MAY-1997; 97US-0047599P.
PR 23-MAY-1997; 97US-0047600P.
PR 23-MAY-1997; 97US-0047601P.
PR 23-MAY-1997; 97US-0047612P.
PR 23-MAY-1997; 97US-0047613P.
PR 23-MAY-1997; 97US-0047614P.
PR 23-MAY-1997; 97US-0047615P.
PR 23-MAY-1997; 97US-0047617P.
PR 23-MAY-1997; 97US-0047618P.
PR 23-MAY-1997; 97US-0047632P.
PR 23-MAY-1997; 97US-0047633P.
PR 06-JUN-1997; 97US-0048964P.
PR 06-JUN-1997; 97US-0048974P.
PR 13-JUN-1997; 97US-0049610P.
PR 08-JUL-1997; 97US-0051926P.
PR 16-JUL-1997; 97US-0052874P.
PR 18-AUG-1997; 97US-0055724P.
PR 22-AUG-1997; 97US-0056630P.
PR 22-AUG-1997; 97US-0056631P.
PR 22-AUG-1997; 97US-0056632P.
PR 22-AUG-1997; 97US-0056636P.
PR 22-AUG-1997; 97US-0056637P.
PR 22-AUG-1997; 97US-0056662P.
PR 22-AUG-1997; 97US-0056664P.
PR 22-AUG-1997; 97US-0056845P.
PR 22-AUG-1997; 97US-0056862P.
PR 22-AUG-1997; 97US-0056864P.
PR 22-AUG-1997; 97US-0056872P.
PR 22-AUG-1997; 97US-0056874P.

PR 22-AUG-1997; 97US-0056875P.
PR 22-AUG-1997; 97US-0056876P.
PR 22-AUG-1997; 97US-0056877P.
PR 22-AUG-1997; 97US-0056878P.
PR 22-AUG-1997; 97US-0056879P.
PR 22-AUG-1997; 97US-0056880P.
PR 22-AUG-1997; 97US-0056881P.
PR 22-AUG-1997; 97US-0056882P.
PR 22-AUG-1997; 97US-0056884P.
PR 22-AUG-1997; 97US-0056886P.
PR 22-AUG-1997; 97US-0056887P.
PR 22-AUG-1997; 97US-0056888P.
PR 22-AUG-1997; 97US-0056889P.
PR 22-AUG-1997; 97US-0056892P.
PR 22-AUG-1997; 97US-0056893P.
PR 22-AUG-1997; 97US-0056894P.
PR 22-AUG-1997; 97US-0056903P.
PR 22-AUG-1997; 97US-0056908P.
PR 22-AUG-1997; 97US-0056909P.
PR 22-AUG-1997; 97US-0056910P.
PR 22-AUG-1997; 97US-0056911P.
PR 05-SEP-1997; 97US-0057650P.
PR 05-SEP-1997; 97US-0057659P.
PR 05-SEP-1997; 97US-0057761P.
PR 12-SEP-1997; 97US-0058785P.
PR 02-OCT-1997; 97US-0061060P.

XX (HUMA-) HUMAN GENOME SCI INC.
PA Ruben SM, Rosen CA, Fischer CL, Soppet DR, Carter KC,
PI Bednarik DP, Endress GA, Yu G, Ni J, Feng P, Young PE, Greene JM;
PI Ferrie AM, Duan R, Hu J, Florence KA, Olsen HS, Ebner R, Brewer LA;
PI Moore PA, Shi Y, Lafleur DW, Li Y, Zeng Z, Kyaw H;
XX WPI; 1998-506364/43.
DR N-ESDB; AAV59572.
XX
XX New isolated human genes and the secreted polypeptide(s) they encode -
PT useful for diagnosis and treatment of e.g. cancers, neurological
PT disorders, immune diseases, inflammation or blood disorders.
XX
PS Claim 1; Page 577; 721pp; English.
XX
XX This sequence represents a secreted human protein encoded by the nucleic
CC acid molecule designated Gene 62 from the human cDNA clone HARP67
CC (deposited as clone ATCC 97900 and ATCC 209046). The gene can be used to
CC generate fusion proteins by linking to the gene to a human immunoglobulin
CC protein as compared to the human protein only. The invention relates to
CC 186 novel genes and their fragments (nucleic acid sequences: AAV59511-
CC V59812; amino acid sequences AAW4731-W75026) which are useful for
CC preventing, treating or ameliorating medical conditions e.g. by protein
CC or gene therapy. Also, pathological conditions can be diagnosed by
CC determining the amount of the new polypeptides in a sample or by
CC determining the presence of mutations in the new polynucleotides.
CC Specific uses are described for each of the 186 polynucleotides, based on
CC which tissues they are most highly expressed in (see AAV59511 for
CC described uses)
XX
SQ Sequence 40 AA;
Query Match 30.9%; Score 30; DB 2; Length 40;
Best Local Similarity 58.3%; Pred. NO. 1.1e+03;
Matches 7; Conservative 1; Mismatches 4; Indels 0; Gaps 0;
OY 3 NMLNSKIAPKIV 14
||| : |||
Db 27 NMLAPRILFFIV 38

RESULT 107
ABG95241
ID ABG95241 standard; protein; 40 AA.

```

XX AC ABG95241;  
 XX DT 15-JAN-2003 (first entry)  
 XX DE Human novel secreted protein #62.  
 XX KW Human; secreted protein; autoimmune disease; chemotaxis;  
 KW rheumatoid arthritis; hyperproliferative disorder; breast neoplasm;  
 KW liver neoplasm cardiovascular disorder; cardiac arrest; skin aging;  
 KW cerebrovascular disorder; cerebral ischemia; angiogenesis; sunburn;  
 KW nervous system disorders; Alzheimer's disease; infection;  
 KW ocular disorder; corneal infection; wound healing; tissue regeneration;  
 KW epithelial cell proliferation; organ transplantation; food additive;  
 KW preservative; nutritional.  
 XX OS Homo sapiens.  
 XX PN US6420526-B1.  
 XX PD 16-JUL-2002.  
 XX PF 08-SEP-1998; 98US-00149476.  
 XX PR 07-MAR-1997; 97US-0038621P.  
 PR 07-MAR-1997; 97US-0040161P.  
 PR 07-MAR-1997; 97US-0040162P.  
 PR 07-MAR-1997; 97US-0040163P.  
 PR 07-MAR-1997; 97US-0040333P.  
 PR 07-MAR-1997; 97US-0040334P.  
 PR 07-MAR-1997; 97US-0040336P.  
 PR 07-MAR-1997; 97US-0040626P.  
 PR 11-APR-1997; 97US-0043311P.  
 PR 11-APR-1997; 97US-0043312P.  
 PR 11-APR-1997; 97US-0043313P.  
 PR 11-APR-1997; 97US-0043314P.  
 PR 11-APR-1997; 97US-0043315P.  
 PR 11-APR-1997; 97US-0043568P.  
 PR 11-APR-1997; 97US-0043569P.  
 PR 11-APR-1997; 97US-0043576P.  
 PR 11-APR-1997; 97US-0043578P.  
 PR 11-APR-1997; 97US-0043580P.  
 PR 11-APR-1997; 97US-0043669P.  
 PR 11-APR-1997; 97US-0043670P.  
 PR 11-APR-1997; 97US-0043671P.  
 PR 11-APR-1997; 97US-0043672P.  
 PR 23-MAY-1997; 97US-0047492P.  
 PR 23-MAY-1997; 97US-0047500P.  
 PR 23-MAY-1997; 97US-0047501P.  
 PR 23-MAY-1997; 97US-0047502P.  
 PR 23-MAY-1997; 97US-0047503P.  
 PR 23-MAY-1997; 97US-0047581P.  
 PR 23-MAY-1997; 97US-0047582P.  
 PR 23-MAY-1997; 97US-0047583P.  
 PR 23-MAY-1997; 97US-0047584P.  
 PR 23-MAY-1997; 97US-0047585P.  
 PR 23-MAY-1997; 97US-0047586P.  
 PR 23-MAY-1997; 97US-0047587P.  
 PR 23-MAY-1997; 97US-0047588P.  
 PR 23-MAY-1997; 97US-0047589P.  
 PR 23-MAY-1997; 97US-0047590P.  
 PR 23-MAY-1997; 97US-0047592P.  
 PR 23-MAY-1997; 97US-0047593P.  
 PR 23-MAY-1997; 97US-0047594P.  
 PR 23-MAY-1997; 97US-0047595P.  
 PR 23-MAY-1997; 97US-0047596P.  
 PR 23-MAY-1997; 97US-0047597P.  
 PR 23-MAY-1997; 97US-0047598P.  
 PR 23-MAY-1997; 97US-0047599P.  
 PR 23-MAY-1997; 97US-0047600P.  
 PR 23-MAY-1997; 97US-0047601P.  
 PR 23-MAY-1997; 97US-0047612P.

PR 23-MAY-1997; 97US-0047613P.  
 PR 23-MAY-1997; 97US-0047614P.  
 PR 23-MAY-1997; 97US-0047615P.  
 PR 23-MAY-1997; 97US-0047617P.  
 PR 23-MAY-1997; 97US-0047618P.  
 PR 23-MAY-1997; 97US-0047632P.  
 PR 23-MAY-1997; 97US-0047633P.  
 PR 06-JUN-1997; 97US-0048964P.  
 PR 06-JUN-1997; 97US-0048974P.  
 PR 13-JUN-1997; 97US-0049610P.  
 PR 08-JUL-1997; 97US-0051926P.  
 PR 16-JUL-1997; 97US-0052874P.  
 PR 18-AUG-1997; 97US-0055724P.  
 PR 22-AUG-1997; 97US-0056630P.  
 PR 22-AUG-1997; 97US-0056631P.  
 PR 22-AUG-1997; 97US-0056632P.  
 PR 22-AUG-1997; 97US-0056636P.  
 PR 22-AUG-1997; 97US-0056637P.  
 PR 22-AUG-1997; 97US-0056662P.  
 PR 22-AUG-1997; 97US-0056664P.  
 PR 22-AUG-1997; 97US-0056845P.  
 PR 22-AUG-1997; 97US-0056862P.  
 PR 22-AUG-1997; 97US-0056864P.  
 PR 22-AUG-1997; 97US-0056872P.  
 PR 22-AUG-1997; 97US-0056874P.  
 PR 22-AUG-1997; 97US-0056875P.  
 PR 22-AUG-1997; 97US-0056876P.  
 PR 22-AUG-1997; 97US-0056877P.  
 PR 22-AUG-1997; 97US-0056878P.  
 PR 22-AUG-1997; 97US-0056879P.  
 PR 22-AUG-1997; 97US-0056880P.  
 PR 22-AUG-1997; 97US-0056881P.  
 PR 22-AUG-1997; 97US-0056882P.  
 PR 22-AUG-1997; 97US-0056884P.  
 PR 22-AUG-1997; 97US-0056886P.  
 PR 22-AUG-1997; 97US-0056887P.  
 PR 22-AUG-1997; 97US-0056888P.  
 PR 22-AUG-1997; 97US-0056889P.  
 PR 22-AUG-1997; 97US-0056892P.  
 PR 22-AUG-1997; 97US-0056893P.  
 PR 22-AUG-1997; 97US-0056894P.  
 PR 22-AUG-1997; 97US-0056903P.  
 PR 22-AUG-1997; 97US-0056908P.  
 PR 22-AUG-1997; 97US-0056909P.  
 PR 22-AUG-1997; 97US-0056910P.  
 PR 22-AUG-1997; 97US-0056911P.  
 PR 05-SEP-1997; 97US-0057650P.  
 PR 05-SEP-1997; 97US-0057669P.  
 PR 05-SEP-1997; 97US-0057761P.  
 PR 12-SEP-1997; 97US-0058785P.  
 PR 02-OCT-1997; 97US-0061060P.  
 PR 06-MAR-1998; 98WO-US004493.  
 XX  
 XX (HUMA-) HUMAN GENOME SCI INC.

Ruben SM, Rosen CA, Fischer CL, Soppet DP, Carter KC;  
 Bednarik DR, Endress GA, Yu G, Ni J, Feng P, Young PE, Greene JM;  
 Ferrie AM, Duan R, Hu J, Florence KA, Olsen HS, Ebner R, Brewer LA;  
 Moore PA, Shi Y, Lafleur DW, Li Y, Zeng Z, Kyaw H;

WPI: 2002-634796/68.  
 N-PSDB; ABS73559.

New isolated human secreted protein for diagnosing, preventing, treating  
 or ameliorating medical conditions and used as a food additive or  
 preservative.

Example 1; SEQ ID NO 381; 129pp; English.

The invention relates to an isolated protein that is one of 186 human  
 secreted proteins, given in the specification, encoded by one of 309 cDNA  
 sequences also given in the specification. The protein is used in a  
 pharmaceutical composition used to prevent, treat or ameliorate a medical

CC condition in e.g. humans, mice, rabbits, goats, horses, cats, dogs,  
CC chickens or sheep. Disorders which are diagnosed or treated include  
CC autoimmune diseases e.g. rheumatoid arthritis, hyperproliferative  
CC disorders e.g. neoplasms of the breast or liver, cardiovascular disorders  
CC e.g. cardiac arrest, cerebrovascular disorders e.g. cerebral ischaemia,  
CC angiogenesis, nervous system disorders e.g. Alzheimer's disease,  
CC infections caused by bacteria, viruses and fungi and ocular disorders  
CC e.g. corneal infection. The polypeptides can also be used to aid wound  
CC healing and epithelial cell proliferation, to prevent skin aging due to  
CC sunburn, to maintain organs before transplantation, for supporting cell  
CC culture of primary tissues, to regenerate tissues and in chemotaxis. The  
CC polypeptides can also be used as a food additive or preservative to  
CC increase or decrease storage capabilities, fat content, lipid, protein,  
CC carbohydrate, vitamins, minerals, cofactors and other nutritional  
CC components. The present sequence represents one of the novel human  
CC secreted proteins of the invention. Note: This sequence did not form part  
CC of the printed specification, but was obtained in electronic format  
CC directly from USPTO at seqdata.uspto.gov/sequence.html?docID=6420526B1  
XX  
SQ Sequence 40 AA;

Query Match 30.9%; Score 30; DB 5; Length 40;  
Best Local Similarity 58.3%; Pred. No. 1.1e+03;  
Matches 7; Conservative 1; Mismatches 4; Indels 0; Gaps 0;  
QY 3 NLLNSKTAFLIV 14  
Dd 27 NLLAFRLIFIV 38  
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||| : |||

RESULT 108  
ABO34435  
ID ABO34435 standard; protein; 40 AA.  
XX  
AC ABO34435;  
XX  
DT 22-SEP-2003 (first entry)  
XX  
DE Region of human secreted protein encoded by cDNA sequence #62.  
XX  
KW Human; secreted protein; hyperproliferative disorder; leukaemia;  
KW breast cancer; wound; reproductive disorder; blood-related disorder;  
KW haemophilia; thrombocytopenia; immunodeficiency; thymic hypoplasia;  
KW Wiskott-Aldrich syndrome; autoimmune disorder; multiple sclerosis;  
KW graft-versus-host disease; Hashimoto's thyroiditis; allergy; asthma;  
KW viral infection; bacterial infection; fungal infection; AIDS; sepsis;  
KW renal disorder; kidney failure; cardiovascular disorder; cytostatic;  
KW angina pectoris; cerebral ischaemia; congenital heart defect;  
KW respiratory disorder; neurological disorder; Alzheimer's disease;  
KW Parkinson's disease; inflammation; Crohn's disease; vulvovaginitis;  
KW immunosuppressive; antibacterial; haemostatic; thrombolytic;  
KW anticoagulant; neuroprotective; thyromimetic; antiallergic;  
KW antiasthmatic; virucide; fungicide; anti-HIV; nephroprotective; antidiabetic;  
KW cerebroprotective; cardiatic; nootropic; antiparkinsonian;  
KW antiinflammatory.  
XX  
OS Homo sapiens.  
XX  
PN US2003049618-A1.  
XX  
PD 13-MAR-2003.  
XX  
PF 16-MAR-2003; 2001US-00809391.  
XX  
PR 07-MAR-1997; 97US-0038621P.  
PR 07-MAR-1997; 97US-0040162P.  
PR 07-MAR-1997; 97US-0040163P.  
PR 07-MAR-1997; 97US-0040333P.  
PR 07-MAR-1997; 97US-0040334P.  
PR 07-MAR-1997; 97US-0040336P.  
PR 07-MAR-1997; 97US-0040626P.  
PR 11-APR-1997; 97US-0043311P.  
PR 11-APR-1997; 97US-0043312P.

PR 11-APR-1997; 97US-0043313P.  
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PR 11-APR-1997; 97US-0043671P.  
PR 11-APR-1997; 97US-0043672P.  
PR 11-APR-1997; 97US-0043674P.  
PR 23-MAY-1997; 97US-0047492P.  
PR 23-MAY-1997; 97US-0047500P.  
PR 23-MAY-1997; 97US-0047502P.  
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PR 23-MAY-1997; 97US-0047593P.  
PR 23-MAY-1997; 97US-0047595P.  
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PR 23-MAY-1997; 97US-0047632P.  
PR 23-MAY-1997; 97US-0047633P.  
PR 06-JUN-1997; 97US-0048964P.  
PR 13-JUN-1997; 97US-0048974P.  
PR 13-JUN-1997; 97US-0049610P.  
PR 08-JUL-1997; 97US-0051926P.  
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PR 18-AUG-1997; 97US-0055724P.  
PR 22-AUG-1997; 97US-0056630P.  
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PR 22-AUG-1997; 97US-0056884P.



22-AUG-1997; 97US-0056886P.  
 22-AUG-1997; 97US-0056887P.  
 22-AUG-1997; 97US-0056888P.  
 22-AUG-1997; 97US-0056889P.  
 22-AUG-1997; 97US-0056890P.  
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 22-AUG-1997; 97US-0056909P.  
 22-AUG-1997; 97US-0056910P.  
 22-AUG-1997; 97US-0056911P.  
 05-SEP-1997; 97US-0057650P.  
 05-SEP-1997; 97US-0057659P.  
 12-SEP-1997; 97US-0057761P.  
 09-OCT-1997; 97US-0058785P.  
 06-MAR-1998; 97US-0061660P.  
 08-SEP-1998; 98WO-US004493.  
 17-MAR-2000; 2000US-0190068P.  
 (RUBE/) RUBEN S M.  
 (ROSE/) ROSEN C A.  
 (SOPP/) SOPPET D R.  
 (CART/) CARTER K C.  
 (BEDN/) BEDNARIK D P.  
 (ENDR/) ENDRESS G A.  
 (YUGG/) YU G.  
 (NIJJ/) NI J.  
 (FENG/) FENG P.  
 (YOUN/) YOUNG P E.  
 (GREE/) GREENE J M.  
 (FERR/) FERRIE A M.  
 (DUAN/) DUAN D R.  
 (HULJ/) HU J.  
 (FLOR/) FLORENCE K A.  
 (OLSE/) OLSEN H S.  
 (FISC/) FISCHER C L.  
 (EBNE/) EBNER R.  
 (BREW/) BREWER L A.  
 (MOOR/) MOORE P A.  
 (SHIY/) SHI Y.  
 (LAFL/) LAFLEUR D W.  
 (LIYY/) LI Y.  
 (ZENG/) ZENG Z.  
 (KYAW/) KYAW H.  
 Ruben SM, Rosen CA, Soppet DR, Carter KC, Bednarik DP, Endress GA, Yu G, Ni J, Feng P, Young PE, Greene JM, Ferrie AM, Duan DR, Hu J, Florence KA, Olsen HS, Fischer CL, Ebner R, Brewer LA, Moore PA, Shi Y, Lafleur DW, Li Y, Zeng Z, Kyaw H; WPI: 2003-521800/49.  
 N-PSDB; AC062702.  
 New genes and its encoded prostate cancer antigen proteins, useful for preventing, treating, ameliorating or diagnosing e.g. prostate cancers, thymic hypoplasia, multiple sclerosis, AIDS, angina pectoris or cerebral ischemia.  
 Claim 3; SEQ ID NO 381; 260pp; English.  
 The present invention relates to the isolation of novel human secreted proteins and the polynucleotide sequences encoding them. The invention also discloses vectors, host cells, antibodies, and recombinant methods for producing human secreted proteins. The polypeptide and polynucleotide sequences for the secreted proteins are useful for preventing, treating, ameliorating or diagnosing medical conditions such as hyperproliferative disorders (e.g. leukaemia or breast cancers), wounds, reproductive disorders, blood-related disorders (e.g. haemophilia or thrombocytopaenia), immunodeficiencies (e.g. Wiskott-Aldrich syndrome or thymic hypoplasia), autoimmune disorders (e.g. graft-versus-host disease, multiple sclerosis or Hashimoto's thyroiditis), allergies (e.g. asthma),

CC viral or bacterial or fungal infections (e.g. AIDS or sepsis), renal disorders (e.g. kidney failure), cardiovascular disorders (e.g. angina pectoris, cerebral ischaemia or congenital heart defects), respiratory disorders, neurological disorders (e.g. Alzheimer's disease or Parkinson's disease), and inflammations (e.g. Crohn's disease). The polynucleotide or polypeptide may also be used as vaccine adjuvants.  
 CC ABO34374-ABO34815 represent human secreted proteins or their fragments.  
 CC Note: The sequence data for this patent did not form part of the printed specification, but was obtained in electronic format directly from the USPTO web site at seqdata.uspto.gov/psipSIDEntry.html  
 CC  
 SQ Sequence 40 AA;  
 Query Match 30.9%; Score 30; DB 6; Length 40;  
 Best Local Similarity 58.3%; Pred. No. 1.1e+03;  
 Matches 7; Conservative 1; Mismatches 4; Indels 0; Gaps 0;  
 QY 3 NHLNSKIAPKIV 14  
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 Db 27 NHLAFRLFFIV 38  
 RESULT 109  
 ADB32063  
 ID ADB32063 standard; peptide; 40 AA.  
 XX  
 AC ADB32063;  
 XX  
 DT 04-DEC-2003 (first entry)  
 XX  
 DE alphaA-integrin alpha subunit alphaA.  
 XX  
 KW Integrin; alphaV-beta3 integrin; RGD peptide ligand; modulator; agonist; antagonist; alphaA.  
 KW  
 XX Unidentified.  
 OS  
 XX WO2003067219-A2.  
 PN  
 XX 14-AUG-2003.  
 PD  
 XX  
 XX 07-FEB-2003; 2003WO-US003903.  
 PF  
 XX  
 XX 07-FEB-2002; 2002US-0354773P.  
 PR  
 XX  
 PA (GEO ) GEN HOSPITAL CORP.  
 XX  
 PI Arnaout AM;  
 XX  
 XX WPI: 2003-663639/62.  
 DR  
 XX  
 PT Screening potential modulators of alphaVbeta3 integrin useful to identify agonists and antagonists uses computer model of three-dimensional structure including a binding site and data from an alphaVbeta3 integrin-ligand complex.  
 PT  
 XX  
 PS Disclosure; Page 5c; 48pp; English.  
 XX  
 CC The invention relates to a method for screening test compounds as potential modulators of alphaV-beta3 integrin using a computer model of the three-dimensional structure of alphaV-beta3 integrin which includes a binding site. The model is based on atomic coordinates of defined alphaV-beta3 integrin amino acids obtained from the structure of a complex of alphaV-beta3 integrin with a known 'RGD peptide' ligand. The method is useful to identify alphaV-beta3 integrin ligands that, because they bind to alphaV-beta3 integrin, may be modulators e.g. agonists or antagonist of alphaV-beta3 integrin activity. It is useful to ascertain whether a specific test compound is a potential modulator and especially to greatly reduce numbers of compounds which must be further tested for their ability to modulate alphaV-beta3 integrin activity. The current sequence represents the alphaA-integrin alpha subunit alphaA.  
 CC  
 XX Sequence 40 AA;  
 SQ

Query Match 30.98; Score 30; DB 7; Length 40;  
Best Local Similarity 46.28; Pred. No. 1.1e+03;  
Matches 6; Conservative 3; Mismatches 4; Indels 0; Gaps 0;

QY 5 LNSKIAPKIVSQE 17  
Db 7 LLSKLRINIISME 19

RESULT 110

ADI23096  
ID ADI23096 standard; protein; 40 AA.

XX AC ADI23096;

DT 22-APR-2004 (first entry)

XX DE Novel human secreted protein seq id 381.

XX KW cytostatic; gene therapy; cancer; human; secreted protein.

XX OS Homo sapiens.

XX PN US2003175858-A1.

XX FD 18-SEP-2003.

XX PF 18-JUN-2001; 2001US-00882171.

XX PR 07-MAR-1997; 97US-0038621P.

PR 07-MAR-1997; 97US-0040162P.

PR 07-MAR-1997; 97US-0040163P.

PR 07-MAR-1997; 97US-0040333P.

PR 07-MAR-1997; 97US-0040334P.

PR 07-MAR-1997; 97US-0040336P.

PR 11-APR-1997; 97US-0043311P.

PR 11-APR-1997; 97US-0043312P.

PR 11-APR-1997; 97US-0043313P.

PR 11-APR-1997; 97US-0043314P.

PR 11-APR-1997; 97US-0043315P.

PR 11-APR-1997; 97US-0043568P.

PR 11-APR-1997; 97US-0043569P.

PR 11-APR-1997; 97US-0043570P.

PR 11-APR-1997; 97US-0043670P.

PR 11-APR-1997; 97US-0043671P.

PR 11-APR-1997; 97US-0043672P.

PR 11-APR-1997; 97US-0043674P.

PR 23-MAY-1997; 97US-0047492P.

PR 23-MAY-1997; 97US-0047500P.

PR 23-MAY-1997; 97US-0047501P.

PR 23-MAY-1997; 97US-0047502P.

PR 23-MAY-1997; 97US-0047503P.

PR 23-MAY-1997; 97US-0047581P.

PR 23-MAY-1997; 97US-0047582P.

PR 23-MAY-1997; 97US-0047583P.

PR 23-MAY-1997; 97US-0047584P.

PR 23-MAY-1997; 97US-0047585P.

PR 23-MAY-1997; 97US-0047586P.

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PR 23-MAY-1997; 97US-0047588P.

PR 23-MAY-1997; 97US-0047589P.

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PR 23-MAY-1997; 97US-0047597P.

PR 23-MAY-1997; 97US-0047598P.  
PR 23-MAY-1997; 97US-0047599P.  
PR 23-MAY-1997; 97US-0047600P.  
PR 23-MAY-1997; 97US-0047601P.  
PR 23-MAY-1997; 97US-0047612P.  
PR 23-MAY-1997; 97US-0047613P.  
PR 23-MAY-1997; 97US-0047614P.  
PR 23-MAY-1997; 97US-0047615P.  
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PR 23-MAY-1997; 97US-0047618P.  
PR 23-MAY-1997; 97US-0047632P.  
PR 23-MAY-1997; 97US-0047633P.  
PR 06-JUN-1997; 97US-0048964P.  
PR 06-JUN-1997; 97US-0048974P.  
PR 13-JUN-1997; 97US-0049610P.  
PR 08-JUL-1997; 97US-0051926P.  
PR 16-JUL-1997; 97US-0052874P.  
PR 18-AUG-1997; 97US-0055724P.  
PR 22-AUG-1997; 97US-0056630P.  
PR 22-AUG-1997; 97US-0056631P.  
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PR 22-AUG-1997; 97US-0056880P.  
PR 22-AUG-1997; 97US-0056881P.  
PR 22-AUG-1997; 97US-0056882P.  
PR 22-AUG-1997; 97US-0056886P.  
PR 22-AUG-1997; 97US-0056887P.  
PR 22-AUG-1997; 97US-0056888P.  
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PR 22-AUG-1997; 97US-0056892P.  
PR 22-AUG-1997; 97US-0056893P.  
PR 22-AUG-1997; 97US-0056894P.  
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PR 22-AUG-1997; 97US-0056911P.  
PR 05-SEP-1997; 97US-0057650P.  
PR 05-SEP-1997; 97US-0057669P.  
PR 12-SEP-1997; 97US-0058785P.  
PR 09-OCT-1997; 97US-0061660P.  
PR 06-MAR-1998; 98WO-US004493.  
PR 08-SEP-1998; 98US-00149476.  
PR 17-MAR-2000; 2000US-0190068P.  
PR 16-MAR-2001; 2001US-00809391.  
XX  
PA (RUBE/) RUBEN S M.  
PA (ROSE/) ROSEN C A.  
PA (SOPP/) SOPPET D R.  
PA (CART/) CARTER K C.  
PA (BEDN/) BEDNARIK D P.  
PA (ENDR/) ENDRESS G A.  
PA (YUGG/) YU G.  
PA (NIJJ/) NI J.  
PA (FENG/) FENG P.  
PA (YOUN/) YOUNG P E.  
PA (GREE/) GREENE J M.  
PA (FERR/) FERRIE A M.

PA (DUAN/) DUAN D R.  
PA (HUJ/) HU J.  
PA (FLOR/) FLORENCE K A.  
PA (OLSE/) OLSEN H S.  
PA (FISC/) FISCHER C L.  
PA (ERNE/) ERNER R.  
PA (BREW/) BREWER L A.  
PA (MOOR/) MOORE P A.  
PA (SHIY/) SHI Y.  
PA (LAF/) LAFLEUR D W.  
PA (LIY/) LI Y.  
PA (ZENG/) ZENG Z.  
PA (KYAW/) KYAW H.  
XX  
PI Ruben SM, Rosen CA, Soppet DR, Carter KC, Bedharik DP;  
PI Andress GA, Yu G, Ni J, Feng P, Young PB, Greene JM, Ferrie AM;  
PI Duan DR, Hu J, Florence KA, Olsen HS, Fischer CL, Ebner R;  
PI Brewer LA, Moore PA, Shi Y, Lafleur DW, Li Y, Zeng Z, Kyaw H;  
XX  
DR WPI: 2003-898535/82.  
DR N-PSDB; ADI22787.  
XX  
PT New nucleic acid molecule, useful for preparing a medicament for  
PT diagnosing, preventing, treating or ameliorating a medical condition  
PT e.g., cancer.  
XX  
PS Claim 11; SEQ ID NO 381; 256pp; English.  
XX  
CC The invention describes an isolated nucleic acid comprising a sequence  
CC having 95 % identity with: a polynucleotide fragment of a sequence not  
CC given in the specification, or its allelic variant; a polynucleotide  
CC fragment of the cDNA sequence; a polynucleotide sequence encoding a  
CC polypeptide, or its fragment, domain, epitope or species homologue; or a  
CC polynucleotide that hybridises under stringent conditions to any one of  
CC the sequences of (a)-(c). The nucleic acid is useful for preparing a  
CC medicament for diagnosing, preventing, treating or ameliorating a medical  
CC condition e.g., cancer. The is the amino acid sequence of a novel human  
CC secreted protein of the invention.  
XX  
SQ Sequence 40 AA;  
XX  
Query Match 30.9%; Score 30; DB 7; Length 40;  
Best Local Similarity 58.3%; Pred. No. 1.le+03;  
Matches 7; Conservative 1; Mismatches 4; Indels 0; Gaps 0;  
OY 3 NHLNKSIAFXIV 14  
||| : |||  
Db 27 NHLAFRLFFIV 38  
XX  
RESULT 111  
ADH74098  
ID ADH74098 standard; protein; 40 AA.  
XX  
AC ADH74098;  
XX  
DT 25-MAR-2004 (first entry)  
XX  
DE Human secreted protein #62.  
XX  
DE human; secreted protein; cancer; haematopoietic disorder;  
KW endocrine disorder; immune system disease; inflammatory disorder.  
XX  
OS Homo sapiens.  
XX  
XX US2003225248-A1.  
XX  
PD 04-DEC-2003.  
XX  
XX 10-JUN-2002; 2002US-00164861.  
PF  
XX  
XX 07-MAR-1997; 97US-0038621P.  
PR  
XX 07-MAR-1997; 97US-0040161P.

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PR 07-MAR-1997; 97US-0040163P.  
PR 07-MAR-1997; 97US-0040333P.  
PR 07-MAR-1997; 97US-0040334P.  
PR 07-MAR-1997; 97US-0040336P.  
PR 07-MAR-1997; 97US-0040626P.  
PR 11-APR-1997; 97US-0043311P.  
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PR 11-APR-1997; 97US-0043314P.  
PR 11-APR-1997; 97US-0043315P.  
PR 11-APR-1997; 97US-0043568P.  
PR 11-APR-1997; 97US-0043569P.  
PR 11-APR-1997; 97US-0043576P.  
PR 11-APR-1997; 97US-0043578P.  
PR 11-APR-1997; 97US-0043580P.  
PR 11-APR-1997; 97US-0043669P.  
PR 11-APR-1997; 97US-0043670P.  
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PR 11-APR-1997; 97US-0043674P.  
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PR 23-MAY-1997; 97US-0047585P.  
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PR 23-MAY-1997; 97US-0047593P.  
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PR 23-MAY-1997; 97US-0047599P.  
PR 23-MAY-1997; 97US-0047600P.  
PR 23-MAY-1997; 97US-0047601P.  
PR 23-MAY-1997; 97US-0047612P.  
PR 23-MAY-1997; 97US-0047613P.  
PR 23-MAY-1997; 97US-0047614P.  
PR 23-MAY-1997; 97US-0047615P.  
PR 23-MAY-1997; 97US-0047617P.  
PR 23-MAY-1997; 97US-0047618P.  
PR 23-MAY-1997; 97US-0047632P.  
PR 23-MAY-1997; 97US-0047633P.  
PR 06-JUN-1997; 97US-0048964P.  
PR 13-JUN-1997; 97US-0048974P.  
PR 08-JUL-1997; 97US-0049610P.  
PR 16-JUL-1997; 97US-0051926P.  
PR 18-AUG-1997; 97US-0052874P.  
PR 22-AUG-1997; 97US-0055724P.  
PR 22-AUG-1997; 97US-0055630P.  
PR 22-AUG-1997; 97US-0055631P.  
PR 22-AUG-1997; 97US-0055632P.  
PR 22-AUG-1997; 97US-0055633P.  
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XX (HUMA-) HUMAN GENOME SCI INC.
XX
PI Rosen CA, Barash SC, Ruben SM;
XX WPI; 2001-483426/52.
DR N-PSDB; AAK58253.
XX
PT Nucleic acids encoding human immune/hematopoietic antigen polypeptides,
PT useful for preventing, diagnosing and/or treating cancers and metastasis.
XX
PS Claim 11; SEQ ID NO 13065; 3071pp + Sequence Listing; English.
XX
CC AAK54951 to AAK64702 encode the human immune/hematopoietic antigen (I)
CC amino acid sequences given in AAM82170 to AAM91921. (I) have cytostatic
CC activity, and can be used in gene therapy and vaccine production. (I)
CC proteins and polynucleotides may be used in the prevention, diagnosis and
CC treatment of diseases associated with inappropriate (I) expression. For
CC example, they may be used to treat disorders associated with decreased
CC expression by rectifying mutations or deletions in a patient's genome
CC that affect the activity of (I) by expressing inactive proteins or to
CC supplement the patients own production of (I). Additionally, (I)
CC polynucleotides may be used to produce the secreted (I), by inserting the
CC nucleic acids into a host cell and culturing the cell to express the
CC protein. (I) proteins and polynucleotides may be used to prevent,
CC diagnose and treat immune/hematopoietic-related diseases, especially
CC cancers and cancer metastases of hematopoietic-derived cells. AAK64703
CC to AAK87694 represent human immune/hematopoietic antigen genomic
CC sequences from the present invention. AAK54942 to AAK54950 and AAM82169
CC represent sequences used in the exemplification of the present invention
XX
SQ Sequence 41 AA;
Query Match 30.9%; Score 30; DB 4; Length 41;
Best Local Similarity 71.4%; Pred. No. 1.2e+03;
Matches 5; Conservative 1; Mismatches 1; Indels 0; Gaps 0;
QY 3 NHELSKI 9
Db ||||| :
4 NHELSL 10
RESULT 113
ABP29414
ID ABP29414 standard; protein; 42 AA.
XX
AC ABP29414;
XX
DT 02-JUL-2002 (first entry)
XX
DE Streptococcus polypeptide SEQ ID NO 8004.
XX
KW Streptococcus; GAS; GBS; group B streptococcus; Streptococcus agalactiae;
KW group A streptococcus; Streptococcus pyogenes; antibacterial;
KW antinflammatory; infection; vaccine; meningitis; gene therapy.
XX
OS Streptococcus pyogenes.
XX
PN WO200234771-A2.
XX

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PD 02-MAY-2002.  
 XX 29-OCT-2001; 2001WO-GB004789.  
 PF  
 XX 27-OCT-2000; 2000GB-00026333.  
 PR 24-NOV-2000; 2000GB-00028727.  
 PR 07-MAR-2001; 2001GB-00005640.  
 XX  
 PA (CHIR-) CHIRON SPA.  
 PA (GENO-) INST GENOMIC RES.  
 XX  
 XX Telford J, Massignani V, Margarit Y Rosi, Grandi G, Fraser C;  
 PI Tettelin H;  
 DR WPI; 2002-352536/38.  
 DR N-ESDB; ABN70045.  
 XX  
 PT New Streptococcus protein for the treatment or prevention of infection or  
 PT disease caused by Streptococcus bacteria, such as meningitis, and for  
 PT detecting a compound that binds to the protein.  
 XX  
 FS Claim 1; Page 3927; 4525pp; English.  
 XX  
 CC The invention relates to a protein (ABP25413-ABP30895) from group B  
 CC streptococcus/GBS (Streptococcus agalactiae) or group A streptococcus/GAS  
 CC (Streptococcus pyogenes), comprising one of 5483 sequences (S1), given in  
 CC the specification. The proteins have antibacterial and antiinflammatory  
 CC activity. (I), nucleic acids encoding (I), ABN66044-ABN71526 and  
 CC antibodies that bind (I) are used in the manufacture of medicaments for  
 CC the treatment or prevention of infection or disease caused by  
 CC Streptococcus bacteria, particularly S. agalactiae and S. pyogenes.  
 CC Nucleic acids encoding (I) are used to detect Streptococcus in a  
 CC biological sample. (I) is used to determine whether a compound binds to  
 CC (I). A composition comprising (I) or a nucleic acid encoding (I), may be  
 CC used as a vaccine or diagnostic composition. The disease caused by  
 CC Streptococcus that is prevented or treated may be meningitis. Nucleic  
 CC acid encoding (I) may be used to recombinantly produce (I) and may be  
 CC used in gene therapy. Antibodies to (I) are used for affinity  
 CC chromatography, immunoassays, and distinguishing/identifying  
 CC Streptococcus proteins  
 XX  
 SQ Sequence 42 AA;  
 Query Match 30.9%; Score 30; DB 5; Length 42;  
 Best Local Similarity 54.5%; Pred No. 1.2e+03;  
 Matches 6; Conservative 3; Mismatches 2; Indels 0; Gaps 0;  
 Qy 5 LNSKIAFKIVS 15  
 Db 6 LNSKESFPFVIS 16  
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 RESULT 114  
 ABB03481  
 ID ABB03481 standard; protein; 46 AA.  
 XX  
 AC ABB03481;  
 XX  
 DT 08-JAN-2002 (first entry)  
 XX  
 DE Human musculoskeletal system related polypeptide SEQ ID NO 1428.  
 XX  
 KW Cytostatic; immunosuppressive; nootropic; neuroprotective; antiviral;  
 KW antiallergic; hepatotropic; antidiabetic; antiinflammatory; antiulcer;  
 KW vulnary; anticonvulsant; antibacterial; antifungal; antiparasitic;  
 KW cardiant; gene therapy; cancer; immune disorder; cardiovascular disorder;  
 KW neurological disease; infection; human; secreted protein;  
 KW musculoskeletal system.  
 OS  
 XX Homo sapiens.  
 XX  
 FN WO200155367-A1.  
 XX

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 XX 17-JAN-2001; 2001WO-US001338.  
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 XX 31-JAN-2000; 2000US-0179065P.  
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 PR 17-NOV-2000; 2000US-0249300P.  
 PR 01-DEC-2000; 2000US-0250160P.  
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 PR 08-DEC-2000; 2000US-0251868P.  
 PR 08-DEC-2000; 2000US-0251869P.  
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 PR 11-DEC-2000; 2000US-0254097P.  
 PR 05-JAN-2001; 2001US-0259678P.  
 (HUMA-) HUMAN GENOME SCI INC.  
 PA Rosen CA, Barash SC, Ruben SM;  
 PI WPI; 2001-451937/48.  
 XX

DR N-PSDB; AAL35063.  
 XX Isolated polypeptide for treating, preventing and/ or prognosing disorders related to the musculoskeletal system including musculoskeletal cancers and also for testing and detection e.g. diagnosis.  
 PT Claim 11; SEQ ID NO 1428; 781pp + Sequence Listing; English.  
 PS The invention relates to novel genes (AAL34669-AAL37666) and proteins (AB03087-AB04109) associated with the musculoskeletal system useful for preventing, treating or ameliorating medical conditions e.g. by protein or gene therapy. The genes are isolated from a range of human tissues disclosed in the specification. The nucleic acids, proteins, antibodies and (ant)agonists are useful in the diagnosis, treatment and prevention of: (a) cancer, e.g. breast and ovarian cancer and other cancers of the adrenal gland, bone, bone marrow, breast, gastrointestinal tract, liver, lung, or urogenital; (b) immune disorders e.g. Addison's disease, allergies, autoimmune haemolytic anaemia, autoimmune thyroiditis, diabetes mellitus; Crohn's disease, multiple sclerosis; rheumatoid arthritis and ulcerative colitis; (c) cardiovascular disorders such as myocardial ischaemias; (d) wound healing; (e) neurological diseases e.g. cerebral anoxia and epilepsy; and (f) infectious diseases such as viral, bacterial, fungal and parasitic infections. Note: The sequence data for this patent did not form part of the printed specification, but was obtained in electronic format directly from WIPO at ftp.wipo.int/pub/published\_pct\_sequences  
 XX SQ Sequence 46 AA;  
 Query Match 30.9%; Score 30; DB 4; Length 46;  
 Best Local Similarity 41.7%; Pred. No. 1.4e+03;  
 Matches 5; Conservative 5; Mismatches 2; Indels 0; Gaps 0;  
 QY 5 LNSKIAPKIVSQ 16  
 Db 28 LSQEVAFKLSQ 39  
 RESULT 115  
 AB012775  
 ID AB012775 standard; protein; 46 AA.  
 XX AC AB012775;  
 XX DT 26-FEB-2003 (first entry)  
 XX DE Novel human musculoskeletal system antigen #395.  
 XX KW Musculoskeletal system antigen; cancer; metastasis; re-vascularisation; thrombosis; arteriosclerosis; mineral content; cardiovascular condition; wound; injury; burn; angiogenesis; ulcer; post-operative tissue repair; limb regeneration; neuronal growth; neurodegenerative disorder;  
 KW Alzheimer's disease; Parkinson's disease; AIDS-related complex;  
 KW chondrocyte growth; bone regeneration; periodontal regeneration;  
 KW tissue transport; bone graft; skin aging; keratinocyte growth; hair loss; melanocyte growth; cell proliferation; cell growth; organ transplant;  
 KW cell differentiation; body height; weight; hair colour; eye colour; skin; percentage of adipose tissue; pigmentation; cosmetic surgery; metabolism;  
 KW biorhythm; cardiac rhythm; depression; tendency for violence; pain;  
 KW reproductive capability; hormone level; endocrine level; appetite;  
 KW libido; memory; stress; storage capability; fat content; lipid content; protein content; carbohydrate content; vitamin content; cofactor content; nutritional component.  
 XX OS Homo sapiens.  
 XX PN US2002147140-A1.  
 XX PD 10-OCT-2002.  
 XX PF 17-JAN-2001; 2001US-00764877.  
 XX PF 31-JAN-2000; 2000US-0179065P.  
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PR 08-DEC-2000; 2000US-0251856P.  
PR 08-DEC-2000; 2000US-0251868P.  
PR 08-DEC-2000; 2000US-0251869P.  
XX (ROSE/) ROSEN C A.  
FA (RUBE/) RUBEN S M.  
FA (BARA/) BARASH S C.  
XX  
PI Rosen CA, Ruben SM, Barash SC;  
XX  
DR WPI; 2003-128199/12.  
DR N-PSDB; AEX58051.  
XX  
PT Isolated nucleic acid molecules encoding musculoskeletal system  
PT associated polypeptides, useful for detecting disorders, e.g. cancer.  
XX  
XX Claim 11; SEQ ID NO 1428; 321pp; English.  
XX  
CC The invention describes an isolated nucleic acid molecule comprising a  
CC sequence encoding musculoskeletal system associated polypeptides useful  
CC for detecting disorders, e.g., cancer or cancer metastases, in animals or  
CC humans. The nucleic acid; stimulates re-vascularization of ischemic  
CC tissues associated with conditions such as thrombosis, arteriosclerosis,  
CC and other cardiovascular conditions; treats wounds due to injuries,  
CC burns, post-operative tissue repair, and ulcers; stimulates angiogenesis  
CC and limb regeneration; stimulates neuronal growth; can treat and prevent  
CC neuronal damage occurring in certain disorders or neurodegenerative

CC conditions, such as, Alzheimer's disease, Parkinson's disease, and AIDS-  
CC related complex; stimulates chondrocyte growth, thus they can be used to  
CC enhance bone and periodontal regeneration and aid in tissue transports or  
CC bone grafts; prevents skin aging due to sunburn by stimulating  
CC keratinocyte growth; prevents hair loss, since FGF family members  
CC activate hair-forming cells and promotes melanocyte growth; stimulates  
CC growth and differentiation of hematopoietic cells and bone marrow cells  
CC when used in combination with other cytokines; maintains organs before  
CC transplantation or for supporting cell culture of primary tissues;  
CC induces tissue of mesodermal origin to differentiate in early embryos;  
CC increases or decreases the differentiation or proliferation of embryonic  
CC stem cells, besides, haematopoietic lineage; modulates mammalian  
CC characteristics, such as, body height, weight, hair colour, eye colour,  
CC skin, percentage of adipose tissue, pigmentation, size, and shape (e.g.,  
CC cosmetic surgery); modulates mammalian metabolism; changes mammal's metal  
CC state or physical state by influencing biorhythms, cardiac rhythms,  
CC depression, tendency for violence, tolerance for pain, reproductive  
CC capabilities, hormonal or endocrine levels, appetite, libido, memory, or  
CC stress; increases or decreases storage capabilities, fat content, lipid,  
CC protein, carbohydrate, vitamins, minerals, cofactors or other nutritional  
CC components. This is the amino acid sequence of a novel human  
CC musculoskeletal system antigen. Note: The sequence data for this patent  
CC did not form part of the printed specification, but was obtained in  
CC electronic format directly from the US patent office at  
CC ftp.seqdata.uspto.gov/sequence.html?DocID=20020147140  
XX  
SQ Sequence 46 AA;  
Query Match 30.9%; Score 30; DB 6; Length 46;  
Best Local Similarity 41.7%; Pred.No. 1.4e+03;  
Matches 5; Conservative 5; Mismatches 2; Indels 0; Gaps 0;  
QY 5 LNSKIAPKIVSQ 16  
Db |::|||::|  
28 LSQEVAFKLSQ 39  
RESULT 116  
ID ADJ28801 standard; protein; 46 AA.  
XX  
AC ADJ28801;  
XX  
DT 20-MAY-2004 (first entry)  
XX  
DE Human musculoskeletal system-associated protein - SEQ ID 1428.  
XX  
KW musculoskeletal system; cytosolic; osteopathic; cancer; osteoporosis;  
XX gene therapy; vaccine; human.  
XX  
OS Homo sapiens.  
XX  
XX  
FN US2004009488-A1.  
XX  
XX 15-JAN-2004.  
XX  
PF 13-SEP-2002; 2002US-00242515.  
XX  
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PR 17-NOV-2000; 2000US-0249211P.  
PR 17-NOV-2000; 2000US-0249212P.  
PR 17-NOV-2000; 2000US-0249213P.  
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PR 17-NOV-2000; 2000US-0249215P.  
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PR 01-DEC-2000; 2000US-0250160P.  
PR 01-DEC-2000; 2000US-0250391P.  
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PR 05-DEC-2000; 2000US-0251988P.  
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PR 08-DEC-2000; 2000US-0251868P.  
PR 08-DEC-2000; 2000US-0251869P.  
PR 08-DEC-2000; 2000US-0251989P.  
PR 08-DEC-2000; 2000US-0251990P.  
PR 11-DEC-2000; 2000US-0254097P.  
PR 05-JAN-2001; 2001US-0259678P.  
PR 17-JAN-2001; 2001US-00764877.  
(HUMA-) HUMAN GENOME SCI INC.  
Rosen CA, Ruben SM, Barash SC;  
WPI: 2004-090458/09.  
N-PSDB; ADJ27778.  
New nucleic acid molecule, useful for preparing a medicament for preventing, treating or ameliorating a medical condition e.g., cancer of musculoskeletal tissues or osteoporosis.  
Claim 11; SEQ ID NO 1428; 289pp; English.  
The invention relates to a novel isolated musculoskeletal system-associated nucleic acid molecule. The nucleic acid of the invention demonstrates cytostatic and osteopathic activities and may be useful for preparing a medicament for preventing, treating or ameliorating a medical condition such as cancer of the musculoskeletal tissues or osteoporosis, possibly via gene therapy or vaccine production. The current sequence is that of the human musculoskeletal system-associated polypeptide of the invention. The current sequence is not shown within the specification per se but is available on the USPTO web-site <http://seqdata.uspto.gov/sequence.html?docID=20040009488>.

XX	Sequence 46 AA;
SQ	Query Match 30.9%; Score 30; DB 8; Length 46; Best Local Similarity 41.7%; Pred. NO. 1.4e+03; Matches 5; Conservative 5; Mismatches 2; Indels 0; Gaps 0;
QY	5 LNSKIAFKIVSQ 16   : :      :   28 LSQEVAFKLSTQ 39
Db	
RESULT 117	
ABP33068	
ID	ABP33068 standard; protein; 50 AA.
AC	ABP33068;
XX	
DT	09-JUL-2002 (first entry)
XX	
DE	Human ORP2041 protein, SEQ ID NO:4082.
XX	
KW	Human; ORP; open reading frame; ORFX; drug screening; diagnosis; disease monitoring; cytokine; cell proliferation; cell differentiation; immune modulation; haematopoiesis regulation; tissue growth; angiogenesis; activin; inhibin; chemotactic; chemokinetic; haemostatic; thrombolytic; tumour inhibition; bodily characteristic; fertility; behaviour; cancer; proliferative disorder; neurological disorder; cardiovascular disease; immune system disorder; organ transplantation; tissue growth disorder; tissue regeneration disorder; diabetes mellitus; hypothyroidism; cholesterol ester storage disease; infection; vulnerability; vasotropic; antipsoriatic; antidiabetic; cytostatic; neutropenic; neuroprotective; antiatherosclerotic; anticoagulant; thrombolytic; cardiant; hypotensive; antithyroid; antiinflammatory; immunomodulator; dermatological; analgesic; virucide; antibacterial; fungicide.
OS	Homo sapiens.
XX	
PN	WO200190366-A2.
XX	
PD	29-NOV-2001.
XX	
EF	24-MAY-2001; 2001WO-US017076.
XX	
PR	24-MAY-2000; 2000US-0206690P.
XX	
PA	(CURA-) CURAGEN CORP.
XX	
PI	Leach MD, Shimkets RA;
XX	
DR	WPI; 2002-106200/14.
N-ESDB; ABN77094.	
XX	
PT	Novel human polypeptides and polynucleotides useful for diagnosing, preventing and treating cardiovascular disease, neurodegenerative, hyperproliferative disorders and disorders related to organ transplantation.
PS	Claim 10; Page 1278; 2508pp; English.
CC	
CC	Sequences ABP31028-ABP35561 represent 4534 novel human proteins designated ORF (open reading frame) 1-4534, and sequences ABN75054- ABN75987 represent cDNAs encoding them. The invention also encompasses polypeptides at least 80% identical to the ORF1-ORF4534 (collectively referred to as ORFX) proteins, polynucleotides at least 85% identical to the ORFX nucleic acid sequences, vectors and host cells comprising ORFX polynucleotides, the recombinant production of ORFX proteins, antibodies specific for ORFX proteins, methods of detecting ORFX polynucleotides and polypeptides, methods of screening for modulators of ORFX expression or activity, and methods of screening individuals for a predisposition to an ORFX-associated disorder. The ORFX proteins of the invention have a wide range of biological activities, such as cytokine, cell proliferation, cell differentiation, immune modulation, haematopoiesis regulation,

XX The invention relates to a protein (ABP25413-ABP30895) from group B  
CC streptococcus/GAS (Streptococcus agalactiae) or group A streptococcus/GAS  
CC (Streptococcus pyogenes), comprising one of 5483 sequences (S1), given in  
CC the specification. The proteins have antibacterial and antiinflammatory  
CC activity. (I), nucleic acids encoding (I), ABN66044-ABN71526 and  
CC antibodies that bind (I) are used in the manufacture of medicaments for  
CC the treatment or prevention of infection or disease caused by  
CC Streptococcus bacteria, particularly S. agalactiae and S. pyogenes.  
CC Nucleic acids encoding (I) are used to detect Streptococcus in a  
CC biological sample. (I) is used to determine whether a compound binds to  
CC Streptococcus comprising (I) or a nucleic acid encoding (I), may be  
CC used as a vaccine or diagnostic composition. The disease caused by  
CC Streptococcus that is prevented or treated may be meningitis. Nucleic  
CC acid encoding (I) may be used to recombinantly produce (I) and may be  
CC used in gene therapy. Antibodies to (I) are used for affinity  
CC chromatography, immunoassays, and distinguishing/identifying  
CC Streptococcus proteins  
XX  
SQ Sequence 32 AA;  
  
Query Match 30.4%; Score 29.5; DB 5; Length 32;  
Best Local Similarity 46.7%; Pred. No. 1.1e+03;  
Matches 7; Conservative 3; Mismatches 4; Indels 1; Gaps 1;  
  
QY 6 NSKIAFKIVSQ-EPA 19  
Db 16 NNRVKIKIACQYEP 30  
  
RESULT 119  
ABP28726  
ID ABP28726 standard; protein; 33 AA.  
XX  
AC ABP28726;  
XX  
DT 02-JUL-2002 (first entry)  
DE Streptococcus polypeptide SEQ ID NO 6628.  
DE Streptococcus; GAS; GBS; group B streptococcus; Streptococcus agalactiae;  
XX group A streptococcus; Streptococcus pyogenes; antibacterial;  
KW antiinflammatory; infection; vaccine; meningitis; gene therapy.  
XX  
OS Streptococcus agalactiae.  
XX  
PN WO200234771-A2.  
XX  
PD 02-MAY-2002.  
XX  
PF 29-OCT-2001; 2001WO-GB004789.  
XX  
PR 27-OCT-2000; 2000GB-00026333.  
PR 24-NOV-2000; 2000GB-00028727.  
PR 07-MAR-2001; 2001GB-00005640.  
XX  
XX {CHIR-) CHIRON SPA.  
PA (CHIR-) INST GENOMIC RES.  
PA  
XX Telford J, Masignani V, Margarit Y RosI, Grandi G, Fraser C;  
PI Tettelin H;  
XX  
XX WPI; 2002-352536/38.  
DR N-PSDB; ABN69357.  
XX  
XX New Streptococcus protein for the treatment or prevention of infection or  
PT disease caused by Streptococcus bacteria, such as meningitis, and for  
PT detecting a compound that binds to the protein.  
XX  
XX Claim 1; Page 3825; 4525pp; English.  
PS  
XX The invention relates to a protein (ABP25413-ABP30895) from group B  
XX streptococcus/GAS (Streptococcus agalactiae) or group A streptococcus/GAS  
CC

CC (Streptococcus pyogenes), comprising one of 5483 sequences (S1), given in  
CC the specification. The proteins have antibacterial and antiinflammatory  
CC activity. (I), nucleic acids encoding (I), ABN66044-ABN71526 and  
CC antibodies that bind (I) are used in the manufacture of medicaments for  
CC the treatment or prevention of infection or disease caused by  
CC Streptococcus bacteria, particularly S. agalactiae and S. pyogenes.  
CC Nucleic acids encoding (I) are used to detect Streptococcus in a  
CC biological sample. (I) is used to determine whether a compound binds to  
CC Streptococcus comprising (I) or a nucleic acid encoding (I), may be  
CC used as a vaccine or diagnostic composition. The disease caused by  
CC Streptococcus that is prevented or treated may be meningitis. Nucleic  
CC acid encoding (I) may be used to recombinantly produce (I) and may be  
CC used in gene therapy. Antibodies to (I) are used for affinity  
CC chromatography, immunoassays, and distinguishing/identifying  
CC Streptococcus proteins  
XX  
SQ Sequence 33 AA;  
  
Query Match 30.4%; Score 29.5; DB 5; Length 33;  
Best Local Similarity 41.2%; Pred. No. 1.1e+03;  
Matches 7; Conservative 4; Mismatches 5; Indels 1; Gaps 1;  
  
QY 1 EPNHLN-SKIAFKIVSQ 16  
Db 7 QPEHINIRIEIMPVSQ 23  
  
RESULT 120  
AAG56269  
ID AAG56269 standard; protein; 44 AA.  
XX  
AC AAG56269;  
XX  
DT 18-OCT-2000 (first entry)  
DE Arabidopsis thaliana protein fragment SEQ ID NO: 72301.  
DE Protein identification; signal transduction pathway; metabolic pathway;  
KW hybridisation assay; genetic mapping; gene expression control; promoter;  
KW termination sequence.  
XX  
OS Arabidopsis thaliana.  
XX  
PN EP1033405-A2.  
XX  
PD 06-SEP-2000.  
XX  
PF 25-FEB-2000; 2000EP-00301439.  
XX  
PR 25-FEB-1999; 99US-0121825P.  
PR 05-MAR-1999; 99US-0123180P.  
PR 09-MAR-1999; 99US-0123548P.  
PR 23-MAR-1999; 99US-0125788P.  
PR 25-MAR-1999; 99US-0126264P.  
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PR 25-AUG-1999; 99US-0150566P.  
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PR 27-AUG-1999; 99US-0151066P.  
PR 27-AUG-1999; 99US-0151080P.  
PR 30-AUG-1999; 99US-0151303P.  
PR 31-AUG-1999; 99US-0151438P.  
PR 01-SEP-1999; 99US-0151930P.  
PR 07-SEP-1999; 99US-0152363P.  
PR 10-SEP-1999; 99US-0153070P.  
PR 13-SEP-1999; 99US-0153758P.  
PR 15-SEP-1999; 99US-0154018P.  
PR 16-SEP-1999; 99US-0154039P.  
PR 20-SEP-1999; 99US-0154779P.  
PR 22-SEP-1999; 99US-0155139P.  
PR 23-SEP-1999; 99US-0155486P.  
PR 24-SEP-1999; 99US-0155659P.  
PR 28-SEP-1999; 99US-0156458P.  
PR 29-SEP-1999; 99US-0156596P.  
PR 04-OCT-1999; 99US-0157117P.  
PR 05-OCT-1999; 99US-0157753P.  
PR 06-OCT-1999; 99US-0157865P.  
PR 07-OCT-1999; 99US-0158029P.  
PR 08-OCT-1999; 99US-0158232P.  
PR 12-OCT-1999; 99US-0158369P.  
PR 13-OCT-1999; 99US-0159283P.  
PR 13-OCT-1999; 99US-0159294P.  
PR 13-OCT-1999; 99US-0159295P.  
PR 14-OCT-1999; 99US-0159329P.  
PR 14-OCT-1999; 99US-0159330P.  
PR 14-OCT-1999; 99US-0159331P.  
PR 14-OCT-1999; 99US-0159637P.  
PR 14-OCT-1999; 99US-0159638P.  
PR 18-OCT-1999; 99US-0159584P.  
PR 21-OCT-1999; 99US-0160741P.  
PR 21-OCT-1999; 99US-0160767P.  
PR 21-OCT-1999; 99US-0160768P.  
PR 21-OCT-1999; 99US-0160770P.  
PR 21-OCT-1999; 99US-0160814P.  
PR 21-OCT-1999; 99US-0160815P.  
PR 22-OCT-1999; 99US-0160980P.  
PR 22-OCT-1999; 99US-0160981P.  
PR 22-OCT-1999; 99US-0160989P.

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PR 25-OCT-1999; 99US-0161404P.
PR 25-OCT-1999; 99US-0161405P.
PR 25-OCT-1999; 99US-0161406P.
PR 26-OCT-1999; 99US-0161359P.
PR 26-OCT-1999; 99US-0161360P.
PR 26-OCT-1999; 99US-0161361P.
PR 28-OCT-1999; 99US-0161920P.
PR 28-OCT-1999; 99US-0161922P.
PR 28-OCT-1999; 99US-0161933P.
PR 29-OCT-1999; 99US-0162142P.

Query Match 30.4%; Score 29.5; DB 3; Length 44;
Best Local Similarity 33.3%; Pred. No. 1.6e+03;
Matches 6; Conservative 5; Mismatches 4; Indels 3; Gaps 1;

QY 2 PNHLNSKIAPKIVSQEPA 19
Db 16 PSHIRSEV---VOPPEPA 30

RESULT 121
AAG60246
ID AAG60246 standard; protein; 44 AA.
XX
AC AAG60246;
XX
DT 18-OCT-2000 (first entry)
DE
DE Arabidopsis thaliana protein fragment SEQ ID NO: 78013.
XX
XX Protein identification; signal transduction pathway; metabolic pathway;
KW hybridisation assay; genetic mapping; gene expression control; promoter;
XX termination sequence.
OS
OS Arabidopsis thaliana.
XX
XX EP1033405-A2.
PN
PD 06-SEP-2000.
XX
XX 25-FEB-2000; 2000EP-00301439.
PF
XX 25-FEB-1999; 99US-0121825P.
PR 05-MAR-1999; 99US-0123180P.
PR 09-MAR-1999; 99US-0123548P.
PR 23-MAR-1999; 99US-0125788P.
PR 25-MAR-1999; 99US-0126264P.
PR 29-MAR-1999; 99US-0126785P.
PR 01-APR-1999; 99US-0127462P.
PR 06-APR-1999; 99US-0128234P.
PR 08-APR-1999; 99US-0128714P.
PR 16-APR-1999; 99US-0129845P.
PR 19-APR-1999; 99US-0130077P.
PR 21-APR-1999; 99US-0130449P.
PR 23-APR-1999; 99US-0130510P.
PR 28-APR-1999; 99US-0130891P.
PR 30-APR-1999; 99US-0131449P.
PR 04-MAY-1999; 99US-0132048P.
PR 04-MAY-1999; 99US-0132407P.
PR 05-MAY-1999; 99US-0132484P.
PR 06-MAY-1999; 99US-0132485P.
PR 06-MAY-1999; 99US-0132486P.
PR 07-MAY-1999; 99US-0132487P.
PR 11-MAY-1999; 99US-0132863P.
PR 14-MAY-1999; 99US-0134256P.
PR 14-MAY-1999; 99US-0134218P.
PR 14-MAY-1999; 99US-0134219P.
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PR 18-MAY-1999; 99US-0134768P.
PR 19-MAY-1999; 99US-0134941P.
PR 20-MAY-1999; 99US-0135124P.
PR 21-MAY-1999; 99US-0135353P.

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PR 24-MAY-1999; 99US-0135629P.
PR 25-MAY-1999; 99US-0136021P.
PR 27-MAY-1999; 99US-0136392P.
PR 28-MAY-1999; 99US-0136782P.
PR 01-JUN-1999; 99US-0137222P.
PR 03-JUN-1999; 99US-0137528P.
PR 04-JUN-1999; 99US-0137502P.
PR 07-JUN-1999; 99US-0137724P.
PR 08-JUN-1999; 99US-0138094P.
PR 10-JUN-1999; 99US-0138540P.
PR 10-JUN-1999; 99US-0138847P.
PR 14-JUN-1999; 99US-0139119P.
PR 16-JUN-1999; 99US-0139452P.
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PR 18-JUN-1999; 99US-0139454P.
PR 18-JUN-1999; 99US-0139455P.
PR 18-JUN-1999; 99US-0139456P.
PR 18-JUN-1999; 99US-0139457P.
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PR 18-JUN-1999; 99US-0139750P.
PR 18-JUN-1999; 99US-0139763P.
PR 21-JUN-1999; 99US-0139817P.
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PR 28-JUN-1999; 99US-0140823P.
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PR 01-JUL-1999; 99US-0141842P.
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PR 02-JUL-1999; 99US-0142055P.
PR 06-JUL-1999; 99US-0142390P.
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PR 12-JUL-1999; 99US-0142977P.
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PR 27-JUL-1999; 99US-0145913P.
PR 27-JUL-1999; 99US-0145918P.
PR 27-JUL-1999; 99US-0145919P.
PR 28-JUL-1999; 99US-0145951P.
PR 02-AUG-1999; 99US-0146386P.
PR 02-AUG-1999; 99US-0146388P.

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PR 02-AUG-1999; 99US-0146389P.
PR 03-AUG-1999; 99US-0147038P.
PR 04-AUG-1999; 99US-0147204P.
PR 04-AUG-1999; 99US-0147302P.
PR 05-AUG-1999; 99US-0147192P.
PR 05-AUG-1999; 99US-0147260P.
PR 06-AUG-1999; 99US-0147303P.
PR 06-AUG-1999; 99US-0147416P.
PR 09-AUG-1999; 99US-0147493P.
PR 09-AUG-1999; 99US-0147935P.
PR 10-AUG-1999; 99US-0148171P.
PR 11-AUG-1999; 99US-0148319P.
PR 12-AUG-1999; 99US-0148341P.
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PR 17-AUG-1999; 99US-0149475P.
PR 18-AUG-1999; 99US-0149426P.
PR 20-AUG-1999; 99US-0149722P.
PR 20-AUG-1999; 99US-0149723P.
PR 20-AUG-1999; 99US-0149829P.
PR 23-AUG-1999; 99US-0149902P.
PR 23-AUG-1999; 99US-0149930P.
PR 25-AUG-1999; 99US-0150566P.
PR 26-AUG-1999; 99US-0150884P.
PR 27-AUG-1999; 99US-0151065P.
PR 27-AUG-1999; 99US-0151066P.
PR 27-AUG-1999; 99US-0151080P.
PR 30-AUG-1999; 99US-0151303P.
PR 31-AUG-1999; 99US-0151438P.
PR 01-SEP-1999; 99US-0151930P.
PR 07-SEP-1999; 99US-0152363P.
PR 10-SEP-1999; 99US-0153070P.
PR 13-SEP-1999; 99US-0153375P.
PR 15-SEP-1999; 99US-0154018P.
PR 16-SEP-1999; 99US-0154039P.
PR 20-SEP-1999; 99US-0154779P.
PR 23-SEP-1999; 99US-0155139P.
PR 23-SEP-1999; 99US-0155486P.
PR 24-SEP-1999; 99US-0155659P.
PR 28-SEP-1999; 99US-0156458P.
PR 28-SEP-1999; 99US-0156596P.
PR 04-OCT-1999; 99US-0157117P.
PR 05-OCT-1999; 99US-0157753P.
PR 06-OCT-1999; 99US-0157865P.
PR 07-OCT-1999; 99US-0158029P.
PR 08-OCT-1999; 99US-0158232P.
PR 12-OCT-1999; 99US-0158369P.
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PR 14-OCT-1999; 99US-0159329P.
PR 14-OCT-1999; 99US-0159330P.
PR 14-OCT-1999; 99US-0159331P.
PR 14-OCT-1999; 99US-0159637P.
PR 14-OCT-1999; 99US-0159638P.
PR 18-OCT-1999; 99US-0159584P.
PR 21-OCT-1999; 99US-0160741P.
PR 21-OCT-1999; 99US-0160767P.
PR 21-OCT-1999; 99US-0160768P.
PR 21-OCT-1999; 99US-0160770P.
PR 21-OCT-1999; 99US-0160814P.
PR 21-OCT-1999; 99US-0160815P.
PR 22-OCT-1999; 99US-0160980P.
PR 22-OCT-1999; 99US-0160981P.
PR 22-OCT-1999; 99US-0160989P.
PR 25-OCT-1999; 99US-0161404P.
PR 25-OCT-1999; 99US-0161405P.
PR 25-OCT-1999; 99US-0161406P.
PR 26-OCT-1999; 99US-0161359P.
PR 26-OCT-1999; 99US-0161360P.
PR 26-OCT-1999; 99US-0161361P.
PR 28-OCT-1999; 99US-0161920P.

PR 28-OCT-1999; 99US-0161992P.
PR 28-OCT-1999; 99US-0161993P.
PR 29-OCT-1999; 99US-0162142P.

Query Match 30.4%; Score 29.5; DB 3; Length 44;
Best Local Similarity 33.3%; Pred. No. 1.6e+03;
Matches 6; Conservative 5; Mismatches 4; Indels 3; Gaps 1;

QY 2 PNLNSKIAFKIVSQEPA 19
Db 16 PSIRSEV---VQPPEPA 30

RESULT 122
AAOI1779
ID AAOI1779 standard; protein; 48 AA.
XX AAOI1779;
XX 06-NOV-2001 (first entry)
XX Human polypeptide SEQ ID NO 25671.
XX Human; cytokine; cell proliferation; cell differentiation; gene therapy;
XX vaccine; peptide therapy; stem cell growth factor; haematopoiesis;
XX tissue growth factor; immunomodulatory; cancer; leukaemia;
XX nervous system disorders; arthritis; inflammation.
XX OS Homo sapiens.
XX WO200164835-A2.
XX 07-SEP-2001.
XX 26-FEB-2001; 2001WO-US004927.
XX 28-FEB-2000; 2000US-00515126.
XX 18-MAY-2000; 2000US-00577409.
XX (HYSE-) HYSEQ INC.
XX Tang YT, Liu C, Drmanac RT;
XX WPI; 2001-514838/56.
XX N-PSDB; AAI91710.
XX Isolated nucleic acids and polypeptides, useful for preventing diagnosing
XX PT and treating e.g. leukemia, inflammation and immune disorders.
XX Claim 20; SEQ ID NO 25671; 1399pp + Sequence Listing; English.
XX The invention relates to human polynucleotides (AAI79941-AAI93841) and
XX the encoded proteins (AAO00010-AAO13910) that exhibit activity relating to
XX cytokine, cell proliferation or cell differentiation or which may induce
XX production of other cytokines in other cell populations. The
XX polynucleotides and polypeptides are useful in gene therapy, vaccines or
XX peptide therapy. The polypeptides have various cytokine-like activities,
XX e.g. stem cell growth factor activity, haematopoiesis regulating
XX activity, tissue growth factor activity, immunomodulatory activity and
XX activin/inhibin activity and may be useful in the diagnosis and/or
XX treatment of cancer, leukaemia, nervous system disorders, arthritis and
XX inflammation. Note: The sequence data for this patent did not form part
XX of the printed specification, but was obtained in electronic format
XX directly from WIPO at ftp.wipo.int/pub/published_pct_sequences
XX SQ Sequence 48 AA;

Query Match 30.4%; Score 29.5; DB 4; Length 48;
Best Local Similarity 37.5%; Pred. No. 1.7e+03;
Matches 6; Conservative 4; Mismatches 3; Indels 3; Gaps 1;

QY 3 PNLNSKIAFKIVSQEPA 18
|||:::|:|
```

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Db          2  NHLSSRVG---VODQP 14

RESULT 123
AAW42197
ID  AAW42197 standard; peptide; 20 AA.
XX
XX  AC  AAW42197;
XX
XX  DT  27-AUG-2003 (revised)
XX  DT  25-MAR-2003 (revised)
XX  DT  16-JUN-1998 (first entry)
XX
XX  DE  T-cell epitope peptide 77 from Japanese cypress pollen antigen Chao2.
XX
XX  KW  Japanese cypress pollen; antigen; T-cell epitope; Chao1; Chao2;
XX  KW  diagnosis; allergy; spring tree pollen disease; pollinosis.
XX
XX  OS  Chamaecyparis obtusa.
XX
XX  FN  WO9747648-A1.
XX
XX  PD  18-DEC-1997.
XX
XX  PF  12-JUN-1997; 97WO-JP02031.
XX
XX  PR  14-JUN-1996; 96JP-00153527.
XX
XX  PA  (MEIP ) MEIJI MILK PROD CO LTD.
XX
XX  PI  Kino K, Dairiri K;
XX
XX  DR  WPI; 1998-052242/05.
XX
XX  PT  T-cell epitope peptide portion of Japanese cypress pollen antigens Chao1
XX  PT  and Chao2 - used for diagnosis and treatment of spring tree pollen
XX  PT  disease.
XX
XX  PS  Claim 2; Page 50; 71pp; Japanese.
XX
XX  CC  The present sequence represents a T-cell epitope peptide from Japanese
XX  CC  cypress pollen antigen Chao2. The present invention describes peptides
XX  CC  which correspond to the T-cell epitope sites on Japanese cypress pollen
XX  CC  antigens Chao1 and Chao2. The peptides can be used as a reagent for the
XX  CC  diagnosis of allergy to Japanese cypress pollen, and as an antigen in the
XX  CC  treatment and prevention of spring tree pollen disease in which the
XX  CC  pollinosis involves reactivity to Japanese cypress pollen. (Updated on 25
XX  CC  -MAR-2003 to correct PI field.) (Updated on 27-AUG-2003 to correct OS
XX  CC  field.)
XX
XX  SQ  Sequence 20 AA;

Query Match          29.9%; Score 29; DB 2; Length 20;
Best Local Similarity 38.5%; Pred. No. 7.4e+02;
Matches 5; Conservative 4; Mismatches 4; Indels 0; Gaps 0;

Qy          7 SKIAFKIVSQEPA 19
           |::|::|::|
Db          6 SNVSLKLTSGKPA 18

RESULT 124
ADB75643
ID  ADB75643 standard; peptide; 20 AA.
XX
XX  AC  ADB75643;
XX
XX  DT  04-DEC-2003 (first entry)
XX
XX  DE  Human TM2 peptide SEQ ID NO:52.
XX
XX  KW  antibody library; CD1 region; CD2 region; VH region; VL region;
XX  KW  immunoglobulin; CD3 region; human; TM2.

Query Match          29.9%; Score 29; DB 2; Length 20;
Best Local Similarity 38.5%; Pred. No. 7.4e+02;
Matches 5; Conservative 4; Mismatches 4; Indels 0; Gaps 0;

Qy          7 SKIAFKIVSQEPA 19
           |::|::|::|
Db          6 SNVSLKLTSGKPA 18

RESULT 125
AAP71423
ID  AAP71423 standard; protein; 21 AA.
XX
XX  AC  AAP71423;
XX
XX  DT  25-MAR-2003 (revised)
XX  DT  03-MAY-1991 (first entry)
XX
XX  DE  Immunomodulator peptide #2 inhibits HIV-T4 interaction.
XX
XX  KW  AIDS; T4 cell receptor; immunomodulation.
XX
XX  OS  Synthetic.
XX
XX  FN  WO8703601-A.
XX
XX  PD  18-JUN-1987.
XX
XX  PF  08-DEC-1986; 86WO-PR000425.
XX
XX  PR  06-DEC-1985; 85FR-00018155.
XX
XX  PA  (INSP ) INST PASTEUR.
XX  PA  (AUPR/) AUFRAY C.
XX  PA  (CNRS ) CENT NAT RECH SCI.
XX
XX

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XX OS Homo sapiens.  
 XX FN WO2003044198-A1.  
 XX PD 30-MAY-2003.  
 XX PF 22-NOV-2002; 2002WO-JP012236.  
 XX PR 22-NOV-2001; 2001JP-00358602.  
 XX PA (UYKE-) UNIV KEIO.  
 XX PI Shimizu N, Takayanagi A, Okui M;  
 XX DR WPI; 2003-449818/42.  
 XX PT Highly stable artificial antibody libraries with super-repository and  
 PT little contamination from unexpressible ones, useful as tool in  
 PT proteomics and e.g. for diagnosis and treating various diseases.  
 XX PS Example 3; Page 34; 108pp; Japanese.  
 XX CC The invention relates to a novel artificial single-stranded antibody  
 CC library with superior-repository. The library is created by using a cDNA  
 CC regions of the VH or VL region of immunoglobulin gene and a fragment  
 CC containing the CD3 region by PCR, respectively, producing VH and VL  
 CC libraries, transferring into a host, and displaying the single-stranded  
 CC antibody on a phage surface. An antibody library of the invention is  
 CC useful as a tool in proteomics and antibody chips and filters, for  
 CC screening ligands for antigens, and for studying protein-DNA interaction,  
 CC diagnosis and treating various diseases. The present sequence represents  
 CC a peptide used in the invention.  
 XX SQ Sequence 20 AA;

Query Match 29.9%; Score 29; DB 7; Length 20;  
 Best Local Similarity 50.0%; Pred. No. 7.4e+02;  
 Matches 5; Conservative 2; Mismatches 3; Indels 0; Gaps 0;  
 Qy 2 PNHLNSKIAP 11  
 |::|::|::|  
 Db 3 PSHLVEKIVY 12  
 |::|::|::|

RESULT 125  
 AAP71423  
 ID AAP71423 standard; protein; 21 AA.  
 XX AC AAP71423;  
 XX DT 25-MAR-2003 (revised)  
 XX DT 03-MAY-1991 (first entry)  
 XX DE Immunomodulator peptide #2 inhibits HIV-T4 interaction.  
 XX KW AIDS; T4 cell receptor; immunomodulation.  
 XX OS Synthetic.  
 XX FN WO8703601-A.  
 XX PD 18-JUN-1987.  
 XX PF 08-DEC-1986; 86WO-PR000425.  
 XX PR 06-DEC-1985; 85FR-00018155.  
 XX PA (INSP ) INST PASTEUR.  
 XX PA (AUPR/) AUFRAY C.  
 XX PA (CNRS ) CENT NAT RECH SCI.

PI	Auffray C, Montagnier L, Klatzmann D, Charron D;
XX	
XX	WPI; 1987-177935/25.
XX	
PT	New peptide derivs. contg. specified exposed tetra:peptide sequences -
PT	inhibiting interaction of AIDS virus with T4 cell receptors.
XX	
PS	Claim 5; Page 48; 57pp; French.
XX	
CC	The peptide is a specific example of a peptide comprising the
CC	tetrapeptide motif RFDS (pref. at position 7 to 10 and optionally having
CC	RE at positions 1 and 2 and/or EL at positions 20 and 21). It interferes
CC	with interaction between the AIDS virus and T4 receptors on lymphocytes.
CC	The peptide also has immunomodulatory activity. It is useful in diagnosis
CC	to detect antibodies to the region of the viral genome containing the
CC	RDS sequence. See also AAP71422 and AAP71424-P71437. (Updated on 25-MAR-
CC	2003 to correct PF field.) (Updated on 25-MAR-2003 to correct PA field.)
XX	
SQ	Sequence 21 AA;
	Query Match 29.9%; Score 29; DB 1; Length 21;
	Best Local Similarity 50.0%; Pred. No. 7.8e+02;
	Matches 6; Conservative 4; Mismatches 2; Indels 0; Gaps 0;
Qy	6 NSKIAFKIVSQE 17
	: : :   ::
Db	9 DSKLAFHHVARE 20
	RESULT 126
AD	ADI40693
ID	ADI40693 standard; peptide; 24 AA.
XX	
AC	ADI40693;
XX	
DT	22-APR-2004 (first entry)
XX	
DE	Nef/SH3 domain inhibitory peptide #6.
XX	
KW	kidney cell dedifferentiation; Nef; Src family tyrosine kinase;
KW	SH3 domain; HIV associated neuropathy; HIVAN; AIDS; dementia; anaemia;
KW	lymphoma; myopathy; cardiomyopathy;
KW	primary HIV-induced disease progression.
XX	
OS	Human immunodeficiency virus 1.
OS	Synthetic.
XX	
Key	Location/Qualifiers
FH	Modified-site 1
FT	/note= "Biotinylated"
XX	
FW	US2003229906-A1.
XX	
PD	11-DEC-2003.
XX	
PF	14-APR-2003; 2003US-00413785.
XX	
PR	15-APR-2002; 2002US-0372557P.
XX	
PA	(GELM/) GELMAN I H.
PA	(KLOT/) KLOTWAN P.
PA	(ZHOU/) ZHOU M M.
XX	
PI	Gelman IH, Klotman P, Zhou MM;
XX	
DR	WPI; 2004-178661/17.
XX	
PT	Inhibiting kidney cell dedifferentiation for treating e.g.; HIV
PT	associated neuropathy by inhibiting the interaction of Nef with a Src
PT	family tyrosine kinase SH3 domain of a polypeptide of the cell.
XX	
PS	Claim 9; SEQ ID NO 6; 42pp; English.
XX	



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QY      3  NHLNSKIAPK 12
      :||| | :
Db      15  DHLNOKTQFE 24

RESULT 128
AAW72756
ID      AAW72756 standard; peptide; 28 AA.
XX
AC      AAW72756;
XX
XX      27-AUG-2003 (revised)
DT      12-JAN-1999 (first entry)
XX
DE      SpeI restriction endonuclease amino terminal 28 residue sequence.
XX
XX      Methylase; SpeI restriction endonuclease; recombinant DNA.
XX
XX      Sphaerotilus.
XX
XX      Key Location/Qualifiers
FT      Misc-difference 20
FT      /note= "unspecified"
XX
XX      EP869174-A2.
XX
XX      07-OCT-1998.
XX
XX      18-MAR-1998; 98EP-00302083.
XX
XX      20-MAR-1997; 97US-00821619.
XX
XX      (NEWE ) NEW ENGLAND BIOLABS INC.
XX
XX      Morgan RD, Chang Z, Mersha FB;
XX
XX      WPI; 1998-508490/44.
XX
XX      New DNA coding for [I SpeI]I restriction endonuclease and methylase -
PT      using host cells premodified with the methylase gene, and using a
PT      purification step which removes non-specific endonuclease and exonuclease
PT      contamination.
XX
XX      Example 1; Page 11; 23pp; English.
XX
XX      The present invention describes isolated DNA from Sphaerotilus species
CC      encoding the SpeI restriction endonuclease. Also described are: (1) a
CC      recombinant DNA vector comprising above DNA; (2) isolated DNA from ATCC
CC      No.98366 encoding the SpeI restriction endonuclease and methylase; (3) a
CC      cloning vector comprising DNA of (2); and (4) host cells transformed by
CC      vectors of (1) or (3). The SpeI enzyme recognises 5'-ACTA/GT-3', and
CC      cells as in (4) can be used to produce recombinant SpeI endonuclease. The
CC      present sequence represents the first 28 residues of SpeI restriction
CC      endonuclease from an example from the present invention. (Updated on 27-
CC      AUG-2003 to correct OS field.)
XX
XX      Sequence 28 AA;
XX
XX      Query Match 29.9%; Score 29; DB 2; Length 28;
XX      Best Local Similarity 55.6%; Pred. No. 1.1e+03;
XX      Matches 5; Conservative 2; Mismatches 2; Indels 0; Gaps 0;
XX
QY      1  EPNHLSKI 9
      :||| | :
Db      4  DPNKLSAL 12

RESULT 129
ABB41309
ID      ABB41309 standard; peptide; 28 AA.
XX
AC      ABB41309;
XX

us-10-799-005a-1.rag
QY      04-FEB-2002 (first entry)
XX
DE      Peptide #8815 encoded by human foetal liver single exon probe.
XX
KW      Human; foetal liver; gene expression; single exon nucleic acid probe.
OS      Homo sapiens.
XX
XX      WO200157277-A2.
XX
XX      09-AUG-2001.
XX
XX      30-JAN-2001; 2001WO-US0000669.
XX
XX      04-FEB-2000; 2000US-0180312P.
XX      26-MAY-2000; 2000US-0207456P.
XX      30-JUN-2000; 2000US-00608408.
XX      03-AUG-2000; 2000US-00632366.
XX      21-SEP-2000; 2000US-0234587P.
XX      27-SEP-2000; 2000US-0236359P.
XX      04-OCT-2000; 2000GB-00024263.
XX
XX      (MOLE-) MOLECULAR DYNAMICS INC.
XX
XX      Penn SG, Hanzel DK, Chen W, Rank DR;
XX
XX      WPI; 2001-483447/52.
XX
XX      Human genome-derived single exon nucleic acid probes useful for analyzing
PT      gene expression in human fetal liver.
XX
XX      Claim 27; SEQ ID NO 33944; 639pp + Sequence Listing; English.
XX
XX      The invention relates to a single exon nucleic acid probe for measuring
CC      human gene expression in a sample derived from human foetal liver. The
CC      single exon nucleic acid probes may be used for predicting, measuring and
CC      displaying gene expression in samples derived from human fetal liver. The
CC      present sequence is a peptide encoded by a single exon nucleic acid probe
CC      of the invention. Note: The sequence data for this patent did not form
CC      part of the printed specification, but was obtained in electronic format
CC      directly from WIPO at ftp.wipo.int/pub/published_pct_sequences
XX
XX      Sequence 28 AA;
XX
XX      Query Match 29.9%; Score 29; DB 4; Length 28;
XX      Best Local Similarity 42.9%; Pred. No. 1.1e+03;
XX      Matches 6; Conservative 3; Mismatches 5; Indels 0; Gaps 0;
XX
QY      5  LNSKIAPKIVSQEP 18
      :||| | :
Db      5  LNSKAGFSIYAFDP 18

RESULT 130
AAM35097
ID      AAM35097 standard; protein; 28 AA.
XX
XX      AAM35097;
XX
XX      17-OCT-2001 (first entry)
XX
DE      Peptide #9134 encoded by probe for measuring placental gene expression.
XX
XX      Probe; microarray; human; placenta; antenatal diagnosis;
XX      genetic disorder.
XX
XX      Homo sapiens.
XX
XX      WO200157272-A2.
XX
XX      09-AUG-2001.
XX
XX      30-JAN-2001; 2001WO-US0000663.
XX

```



CC samples, which may enable the improved diagnosis and treatment of cancers  
 CC such as lymphoma, leukaemia and myeloma. The present sequence is a  
 CC protein encoded by one of the probes of the invention

XX SQ Sequence 28 AA;

Query Match 29.9%; Score 29; DB 4; Length 28;  
 Best Local Similarity 42.9%; Pred. No. 1.1e+03;  
 Matches 6; Conservative 3; Mismatches 5; Indels 0; Gaps 0;

QY 5 LNSKIAPKIVSQEP 18  
 |||:|:|:  
 Db 5 LKSKASFSIYAFDP 18

RESULT 133

NAME2177  
 ID AAM62177 standard; protein; 28 AA.

XX AC

XX AAM62177;

XX 05-NOV-2001 (first entry)

XX Human brain expressed single exon probe encoded protein SEQ ID NO: 34282.

XX Human; brain expressed exon; gene expression analysis; probe: microarray;  
 KW Alzheimer's disease; multiple sclerosis; schizophrenia; epilepsy; cancer.

XX OS Homo sapiens.

XX WO200157275-A2.

XX 09-AUG-2001.

XX 30-JAN-2001; 2001WO-US000667.

XX 04-FEB-2000; 2000US-0180312P.

XX 26-MAY-2000; 2000US-0207456P.

XX 30-JUN-2000; 2000US-00608408.

XX 03-AUG-2000; 2000US-00632366.

XX 21-SEP-2000; 2000US-0234687P.

XX 27-SEP-2000; 2000US-0236359P.

XX 04-OCT-2000; 2000GB-00024263.

XX (MOLE-) MOLECULAR DYNAMICS INC.

XX Penn SG, Hanzel DK, Chen W, Rank DR;

XX WPI; 2001-483446/52.

XX Single exon nucleic acid probes for analyzing gene expression in human  
 PT brains.

XX Example 4; SEQ ID NO 34282; 650pp + Sequence Listing; English.  
 PS The present invention provides a number of single exon nucleic acid  
 CC probes which are derived from genomic sequences expressed in the human  
 CC brain. They can be used to measure gene expression in brain cell samples,  
 CC which may enable the diagnosis and improved treatment of nervous system  
 CC diseases such as Alzheimer's disease, multiple sclerosis, schizophrenia,  
 CC epilepsy and cancers. The present sequence is a protein encoded by one of  
 CC the probes of the invention

XX SQ Sequence 28 AA;

Query Match 29.9%; Score 29; DB 4; Length 28;  
 Best Local Similarity 42.9%; Pred. No. 1.1e+03;  
 Matches 6; Conservative 3; Mismatches 5; Indels 0; Gaps 0;

QY 5 LNSKIAPKIVSQEP 18

|||:|:|:  
 Db 5 LKSKASFSIYAFDP 18

RESULT 134

ABG56752

ID ABG56752 standard; peptide; 28 AA.

XX AC

XX ABG56752;

XX 25-FEB-2003 (first entry)

XX Human liver peptide, SEQ ID No 35400.

XX Human; liver; cirrhosis; hyperlipoproteinaemia; hyperlipidaemia;  
 KW hypercholesterolaemia; coronary heart disease.

XX OS Homo sapiens.

XX WO200157273-A2.

XX 09-AUG-2001.

XX 30-JAN-2001; 2001WO-US000664.

XX 04-FEB-2000; 2000US-0180312P.

XX 26-MAY-2000; 2000US-0207456P.

XX 30-JUN-2000; 2000US-00608408.

XX 03-AUG-2000; 2000US-00632366.

XX 21-SEP-2000; 2000US-0234687P.

XX 27-SEP-2000; 2000US-0236359P.

XX 04-OCT-2000; 2000GB-00024263.

XX (MOLE-) MOLECULAR DYNAMICS INC.

XX Penn SG, Hanzel DK, Chen W, Rank DR;

XX WPI; 2001-488898/53.

XX Human genome-derived single exon nucleic acid probes useful for analyzing  
 PT gene expression in human adult liver.

XX Claim 27; SEQ ID NO 35400; 650pp; English.

XX The invention relates to a single exon nucleic acid probe (SENP) (I) for  
 CC measuring human gene expression in a sample derived from human adult  
 CC liver, comprising one of 13109 defined nucleotide sequences given in the  
 CC specification (or complements/ fragments). The probe hybridises at high  
 CC stringency to a nucleic acid molecule expressed in the human adult liver.  
 CC (I) may be used for predicting, measuring and displaying gene expression  
 CC in samples derived from human adult liver. The genes identified may be  
 CC involved in genetic liver diseases such as cirrhosis,  
 CC hyperlipoproteinaemia, hyperlipidaemia and hypercholesterolaemia which is  
 CC associated with coronary heart disease. ABG47348-ABG59930 represent human  
 CC liver single exon encoded peptides of the invention. Note: The sequence  
 CC information for this patent does not appear in the printed specification  
 CC but was obtained in electronic format directly from WIPO at  
 CC ftp.wipo.int/pub/published\_pct\_sequences

XX SQ Sequence 28 AA;

Query Match 29.9%; Score 29; DB 4; Length 28;

Best Local Similarity 42.9%; Pred. No. 1.1e+03;

Matches 6; Conservative 3; Mismatches 5; Indels 0; Gaps 0;

QY 5 LNSKIAPKIVSQEP 18

|||:|:|:  
 Db 5 LKSKASFSIYAFDP 18

RESULT 135

ABG44713

ID ABG44713 standard; peptide; 28 AA.

XX AC

XX ABG44713;

DT 19-AUG-2002 (first entry)  
DE Human peptide encoded by genome-derived single exon probe SEQ ID 34378.  
XX  
XX Human; single exon probe; asthma; lung cancer; COPD; ILD;  
KW chronic obstructive pulmonary disease; interstitial lung disease;  
KW familial idiopathic pulmonary fibrosis; neurofibromatosis;  
KW tuberous sclerosis; Gaucher's disease; Niemann-Pick disease;  
KW Hermansky-Pudlak syndrome; sarcoidosis; pulmonary haemosiderosis;  
KW pulmonary histiocytosis; lymphangioleiomyomatosis; Karagener syndrome;  
KW pulmonary alveolar proteinosis; fibrocystic pulmonary dysplasia;  
KW primary ciliary dyskinesia; pulmonary hypertension;  
KW hyaline membrane disease.  
XX  
OS Homo sapiens.  
XX WO200186003-A2.  
XX 15-NOV-2001.  
XX 30-JAN-2001; 2001WO-US000665.  
XX 04-FEB-2000; 2000US-0180312P.  
XX 26-MAY-2000; 2000US-0207456P.  
XX 30-JUN-2000; 2000US-00608408.  
XX 03-AUG-2000; 2000US-00632366.  
XX 21-SEP-2000; 2000US-0234687P.  
XX 27-SEP-2000; 2000US-0236359P.  
XX 04-OCT-2000; 2000GB-00024263.  
XX (MOLE-) MOLECULAR DYNAMICS INC.  
XX Penn SG, Hanzel DK, Chen W, Rank DR;  
PI WPI; 2002-114183/15.  
XX Spatially-addressable set of single exon nucleic acid probes, used to  
XX measure gene expression in human lung samples.  
XX Claim 27; SEQ ID NO 34378; 634pp; English.  
XX The invention relates to a spatially-addressable set of single exon  
XX nucleic acid probes for measuring gene expression in a sample derived  
XX from human lung comprising single exon nucleic acid probes having one of  
XX 12614 nucleic acid sequences mentioned in the specification, or their  
XX complements or the 12387 open reading frames derived from the 12614  
XX probes. Also included are a microarray comprising the novel set of probes  
XX; the novel set of probes which hybridise at high stringency to a nucleic  
XX acid expressed in the human lung; measuring gene expression in a sample  
XX derived from human lung, comprising (a) contacting the array with a  
XX collection of detectably labeled nucleic acids derived from human lung  
XX mRNA, and (b) measuring the label detectably bound to each probe of the  
XX array; identifying exons in a eukaryotic genome, comprising (a)  
XX algorithmically predicting at least one exon from genomic sequences of  
XX the eukaryote; and (b) detecting specific hybridisation of detectably  
XX labeled nucleic acids from eukaryote lung mRNA, to a single exon probe,  
XX having a fragment identical to the predicted exon, the probe is included  
XX in the above mentioned microarray; assigning exons to a single gene,  
XX comprising (a) identifying exons from genomic sequence by the method  
XX above and (b) measuring the expression of each of the exons in several  
XX tissues and/or cell types using hybridisation to a single exon  
XX microarrays having a probe with the exon, where a common pattern of  
XX expression of the exons in the tissues and/or cell types indicates that  
XX the exons should be assigned to a single gene; a peptide comprising one  
XX of 12011 sequences, mentioned in the specification, or encoded by the  
XX probes/open reading frames (ORF). The probes are used for gene expression  
XX analysis, and for identifying exons in a gene, particularly using human  
XX lung derived mRNA and for the study of lung diseases such as asthma, lung  
XX cancer, chronic obstructive pulmonary disease (COPD), interstitial lung  
XX disease (ILD), familial idiopathic pulmonary fibrosis, neurofibromatosis,  
XX tuberous sclerosis, Gaucher's disease, Niemann-Pick disease, Hermansky-  
XX Pudlak syndrome, sarcoidosis, pulmonary haemosiderosis, pulmonary  
XX histiocytosis, lymphangioleiomyomatosis, pulmonary alveolar proteinosis,

CC Karagener syndrome, fibrocystic pulmonary dysplasia, primary ciliary  
CC dyskinesia, pulmonary hypertension and hyaline membrane disease. The  
CC present sequence is a peptide/protein encoded by a single exon probe of  
CC the invention. Note: the sequence data for this patent did not form part  
CC of the printed specification, but was obtained in electronic format  
CC directly from WIPO at ftp.wipo.int/pub/published\_pct\_sequences  
XX  
XX Sequence 28 AA;  
XX  
XX Query Match 29.9%; Score 29; DB 5; Length 28;  
XX Best Local Similarity 42.9%; Pred. No. 1.e+03;  
XX Matches 6; Conservative 3; Mismatches 5; Indels 0; Gaps 0;  
Qy 5 LNSKIAPKIVSQEP 18  
Db 5 LKSKASFSIYAFDP 18  
RESULT 136  
ABG93608  
ID ABG93608 standard; peptide; 28 AA.  
XX AC ABG93608;  
XX 25-NOV-2002 (first entry)  
XX Human P-glycoprotein tryptic peptide #128.  
XX Human; P-glycoprotein; tryptic digest; proteolytic cleavage product;  
XX diabetes; Parkinson's disease; Alzheimer's disease; malaria; cholera;  
XX human immunodeficiency virus infection; influenza; rabies; diphtheria;  
XX cancer; multi-drug resistance; MDR.  
XX OS Homo sapiens.  
XX EP1223534-A1.  
XX 17-JUL-2002.  
XX 11-JAN-2002; 2002EP-00075095.  
XX 14-JAN-2001; 2001IL-00140881.  
XX 19-OCT-2001; 2001US-00982172.  
XX (KATZ/) KATZ E I.  
XX Katz EI;  
XX WPI; 2002-645691/70.  
XX Generating amino acid sequences representative of desired polypeptide, by  
XX computationally generating proteolytic cleavage products, analyzing and  
XX selecting the set of products, thus generating amino acid sequences.  
XX Example 1; Page 15; 124pp; English.  
XX The invention relates to generating set of amino acid sequences (AAS)  
XX representative of one desired polypeptide (I), involving computationally  
XX generating a number of proteolytic cleavage products (PCP) from (I),  
XX analysing the PCP according to one parameter defining a characteristic of  
XX AAS and selecting a set of PCP according to a preset criteria for each  
XX parameter, thus generating the set of AAS representative of (I). Also  
XX included are (1) a computer readable storage media (II) comprising a  
XX database of amino acid sequences corresponding to the polypeptide of  
XX interest; (2) a system (III) for generating a database of amino acid  
XX sequences corresponding to a polypeptide of interest, comprises a  
XX processing unit which executes a software application configured for  
XX generating the number of proteolytic cleavage products from one  
XX polypeptide of interest, and analysing the number of proteolytic cleavage  
XX products according to one parameter defining a characteristic of amino  
XX acid sequence; (3) a kit for quantifying at least one polypeptide of  
XX interest, comprises a number of peptides or antibodies each capable of  
XX specifically recognising at least one peptide, where the number of

CC peptides is generated according to information derived from computational  
 CC analysis of the polypeptide of interest; and (4) quantifying one  
 CC polypeptide of interest in a biological sample, involving contacting the  
 CC biological sample with proteolytic agent, so as to obtain a proteolysed  
 CC biological sample, contacting the proteolysed biological sample with at  
 CC least one antibody and at least one peptide of a number of peptides, and  
 CC detecting presence, absence and/or level of antibody binding to thus  
 CC quantify one polypeptide of interest in the biological sample. The method  
 CC is useful for generating at least one antibody specific to a polypeptide  
 CC of interest. The peptides or antibodies generated may be used to diagnose  
 CC diabetes, Parkinson's disease, Alzheimer's disease, human  
 CC immunodeficiency virus infection, malaria, cholera, influenza, rabies,  
 CC diphtheria, cancer (e.g. breast, colon, cervix, melanoma, lung, ovary,  
 CC pancreas, prostate, lymphomas and leukaemias). The present sequence is a  
 CC predicted tryptic peptide from human P-glycoprotein generated to form  
 CC part of a kit for identifying multi-drug (MDR) resistance associated  
 CC proteins

XX  
 SQ Sequence 28 AA;

Query Match 29.9%; Score 29; DB 5; Length 28;  
 Best Local Similarity 100.0%; Pred. No. 1.1e+03;  
 Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 13 IVSQEP 18  
 : |||||  
 Db 5 IVSQEP 10

RESULT 137

AAAB27777  
 ID AAB27777 standard; protein; 29 AA.

AC AAB27777;

XX AAB27777;

DT 30-JAN-2001 (first entry)

XX Sequence homologous to protein fragment encoded by gene 42.

XX Cytostatic; immunosuppressive; nootropic; neuroprotective; antiviral;  
 KW antiallergic; hepatotropic; antidiabetic; antiinflammatory; antitumor;  
 KW vulnerary; anticonvulsant; antibacterial; antifungal; antiparasitic;  
 KW cardiant; gene therapy; cancer; immune disorder; cardiovascular disorder;  
 KW neurological disease; infection; human; secreted protein.

XX Homo sapiens.

XX WO200055201-A1.

XX 21-SEP-2000.

XX 09-MAR-2000; 2000WO-US006059.

XX 12-MAR-1999; 99US-0124096P.

XX 03-DEC-1999; 99US-0168622P.

XX (HUMA-) HUMAN GENOME SCI INC.

XX Rosen CA, Ruben SM, Komatsoulis G;

XX WPI; 2000-628182/60.

XX Novel human secreted proteins useful for diagnosis, prevention and  
 PT treatment of disorders including neurological, cell proliferative,  
 PT cardiovascular, autoimmune/inflammatory disorders and microbial  
 PT infections.

XX Disclosure; Page 54; 427pp; English.

XX The invention relates to the isolation of genes AAC59157-C59205 encoding  
 CC the human secreted proteins AAB27682-B27730. This sequence represents a  
 CC peptide fragment homologous to the protein encoded by the gene given in  
 CC the descriptor line. The sequence is a search result from a BLASTX

CC homology search. The genes and proteins are useful for preventing,  
 CC ameliorating or treating medical conditions, e.g. by protein or gene  
 CC therapy. The genes are isolated from a range of human tissues disclosed  
 CC in the specification. The nucleic acids, proteins, antibodies and  
 CC (ant)agonists are useful in the diagnosis, treatment and prevention of:  
 CC (a) cancer, e.g. breast and ovarian cancer, and other cancers of the  
 CC adrenal gland, bone, bone marrow, breast, gastrointestinal tract, liver,  
 CC lung, or urogenital; (b) immune disorders e.g. Addison's disease,  
 CC allergies, autoimmune haemolytic anaemia, autoimmune thyroiditis,  
 CC diabetes mellitus, Crohn's disease, multiple sclerosis, rheumatoid  
 CC arthritis and ulcerative colitis; (c) cardiovascular disorders such as  
 CC myocardial ischaemias; (d) wound healing; (e) neurological diseases e.g.  
 CC cerebral anoxia and epilepsy; and (f) infectious diseases such as viral,  
 CC bacterial, fungal and parasitic infections

XX SQ Sequence 29 AA;

Query Match 29.9%; Score 29; DB 3; Length 29;  
 Best Local Similarity 71.4%; Pred. No. 1.2e+03;  
 Matches 5; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

QY 1 EPNELNS 7

: |||||

Db 6 QENELNS 12

RESULT 138

AAW19976

ID AAW19976 standard; peptide; 31 AA.

XX AAW19976;

XX 25-MAR-2003 (revised)

DT 18-AUG-1997 (first entry)

XX Fibronectin fragment P2, corresponds to aa 625-655.

XX Fibronectin; structural; promote; disulphide bridge; cell culture;  
 KW extracellular matrix; migration; growth; metastasis; tumour; cancer;  
 KW neoplasia; fibronectin-fibronectin binding site; self-assembly;  
 KW type III repeat; C-terminal type I repeat.

OS Homo sapiens.

XX US5629291-A.

XX 13-MAY-1997.

XX 17-NOV-1994; 94US-00340812.

XX 31-JAN-1992; 92US-00829462.

PR 16-FEB-1993; 93US-00021626.

XX (LJOL-) LA JOLLA CANCER RES FOUND.

XX Morla A, Ruoslahti EI;

XX WPI; 1997-280300/25.

XX Promoting extracellular fibronectin matrix formation - by contacting  
 PT cells with fibronectin fragments, useful in, e.g. cell culture.

XX Example 1; Col 11; 42pp; English.

XX AAW19976 and AAW19977 are peptide fragments of human fibronectin (FN)  
 CC type III-I repeat region. Certain peptides derived from this region  
 CC promote extracellular FN matrix formation in a cellular system in the  
 CC presence of FN. The peptides promote FN self-assembly by forming  
 CC disulphide cross-linked FN. Normal fibroblasts in tissue culture secrete  
 CC FN and assemble it into a matrix that is essential for adhesion, growth  
 CC and migration. Tumour cells fail to assemble FN into the extracellular  
 CC matrix (ECM), the lack of ECM is thought to contribute to the invasive  
 CC properties of tumour cells. Other peptides derived from the C-terminal

CC type I repeat of FN are used to inhibit matrix formation by blocking the  
 CC FN-FN, self assembly binding site. The peptides are used to control  
 CC biological processes related to extracellular FN matrix formation, e.g in  
 CC cell culture, directing tissue regeneration and ameliorating certain  
 CC pathological conditions. Matrix formation- inhibiting peptides can  
 CC prevent scar formation. (Updated on 25-MAR-2003 to correct PF field.)  
 XX  
 SQ Sequence 31 AA;

Query Match 29.9%; Score 29; DB 2; Length 31;  
 Best Local Similarity 83.3%; Pred. NO. 1.3e+03;  
 Matches 5; Conservative 0; Mismatches 1; Indels 0; Gaps 0;

Qy 2 PNHLNS 7  
 | ||||  
 Db 5 PGLHNS 10

RESULT 139  
 AAW82987  
 ID AAW82987 standard; peptide; 31 AA.

AC AAW82987;  
 XX  
 DT 10-FEB-1999 (first entry)  
 XX  
 DE Human fibronectin III-1 repeat peptide fragment #2.

XX Human; fibronectin; type III repeat; binding site; inhibition;  
 KW tumour cell migration; extracellular matrix assembly; scar formation;  
 KW implant.

XX Homo sapiens.

OS US5837813-A.

PN 17-NOV-1998.

PD 01-JUN-1995; 95US-00460421.

PF 31-JAN-1992; 92US-00829462.

PR 16-FEB-1993; 93US-00021626.

PR 17-NOV-1994; 94US-00340812.

XX (LOOL-) LA JOLLA CANCER RES FOUND.

PI Ruoslahti EI, Morla A;

XX WPI; 1999-023534/02.

XX New recombinant fragments of the III-1 repeat of fibronectin - contain

PT fibronectin binding site so modulate extracellular matrix assembly and

PT cell migration and increase cell binding to surfaces.

XX Example 1; Col 11; 45pp; English.

XX The present sequence represents a fragment of the III-1 repeat of

CC fibronectin. The present invention describes a fragment of the III-1

CC repeat of fibronectin which is capable of binding fibronectin. The

CC protein fragment inhibits fibronectin-fibronectin binding, and so

CC modulates (enhances or inhibits, depending on concentration) fibronectin

CC extracellular matrix assembly and related processes. The protein fragment

CC can be used to inhibit the migration of tumour cells, inhibit scar

CC formation and promote cell attachment to surfaces such as implants. The

CC protein fragment may also be used to target molecules to fibronectin-

CC containing tissues and cells, or for affinity isolation of fibronectin

Qy 2 PNHLNS 7  
 | ||||  
 Db 5 PGLHNS 10

RESULT 140  
 AAB39245

ID AAB39245 standard; protein; 31 AA.

XX AAB39245;

DT 02-FEB-2001 (first entry)

DE Gene 9 human secreted protein homologous amino acid sequence #125.

XX Human; secreted protein; immunosuppressive; antiarthritic; antirheumatic;  
 KW antiproliferative; cytostatic; cardiant; vasotropic; cerebroprotective;  
 KW nootropic; neuroprotective; antibacterial; virucide; fungicide; neoplasm;  
 KW ophthalmological; autoimmune disease; rheumatoid arthritis; angiogenesis;  
 KW hyperproliferative disorder; cardiovascular disorder; infection;  
 KW cerebrovascular disorder; nervous system disorder; ocular disorder;  
 KW wound healing; chemotaxis.

XX Homo sapiens.

OS WO200056754-A1.

PN 28-SEP-2000.

PD 16-MAR-2000; 2000WO-US006792.

PF 19-MAR-1999; 99US-0125362P.

PR 10-DEC-1999; 99US-0169980P.

XX (HUMA-) HUMAN GENOME SCI INC.

XX Rosen GA, Ruben SM, Komatsoulis G;

XX WPI; 2000-579483/54.

XX Isolated nucleic acid molecule encoding a human secreted protein is used

PT in preventing, treating or ameliorating a medical condition.

XX Disclosure; Page 25; 434pp; English.

XX The polynucleotide sequences given in AAC74223-C74279 encode the human  
 CC secreted proteins represented in AAB39179-B39226. Sequences AAB39227-  
 CC B39308 are alternative proteins encoded by the genes, and also protein  
 CC sequences with which they share homology. The proteins have activities  
 CC based on the tissues and cells in which they are expressed. Examples of  
 CC activities include: immunosuppressive; antiarthritic; antirheumatic;  
 CC antiproliferative; cytostatic; cardiant; vasotropic; cerebroprotective;  
 CC nootropic; neuroprotective; antibacterial; virucide; fungicide; and  
 CC ophthalmological. The human secreted proteins, polynucleotides,  
 CC antagonists and agonists of the invention may be useful in the treatment,  
 CC prevention, and/or diagnosis of various disease, disorders and conditions  
 CC such as autoimmune diseases e.g. rheumatoid arthritis, hyperproliferative  
 CC disorders e.g. neoplasms of the breast or liver, cardiovascular disorders  
 CC e.g. cardiac arrest, cerebrovascular disorders e.g. cerebral ischaemia,  
 CC angiogenesis, nervous system disorders e.g. Alzheimer's disease,  
 CC infections caused by bacteria, viruses and fungi and ocular disorders  
 CC e.g. corneal infection. The polypeptides can also be used to aid wound  
 CC healing and epithelial cell proliferation, to regenerate tissues,  
 CC maintain organs before transplantation, in chemotaxis and as a food  
 CC additive or preservative e.g. to increase storage capabilities. Sequences  
 CC AAC74214-C74222 and AAB39178 are used during the isolation and  
 CC characterisation of the genes of the invention

XX Sequence 31 AA;

Query Match 29.9%; Score 29; DB 3; Length 31;

Best Local Similarity 71.4%; Pred. No. 1.3e+03;

Matches 5; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

QY 1 EPNHLS 7  
 Db : ||||| 7  
 1 QENHLS 7

RESULT 141  
 ABO55532  
 ID ABO55532 standard; protein; 32 AA.  
 XX AC ABO55532;  
 XX DT 29-JUL-2004 (first entry)  
 XX DE Human genome derived single exon protein #1766.  
 XX Human; gene expression; single exon probe; microarray;  
 KW alternative splicing event; genomic alteration.  
 XX OS Homo sapiens.  
 XX PN US2003194704-A1.  
 XX PD 16-OCT-2003.  
 XX PF 03-APR-2002; 2002US-00029386.  
 XX PR 03-APR-2002; 2002US-00029386.  
 XX PA (PENN/) PENN S G.  
 XX PA (RANK/) RANK D R.  
 XX PA (HANZ/) HANZEL D K.  
 XX PI Penn SG, Rank DR, Hanzel DK;  
 XX WPI: 2004-119264/12.

PT New human genome-derived single exon nucleic acid probes useful for human  
 gene expression analysis, for identifying or characterizing alternative  
 splicing events, for assessing genomic alterations or as tools for  
 surveying tissues.

XX Claim 45; SEQ ID NO 29166; 80pp; English.

XX The invention relates to a nucleic acid probe for measuring human gene  
 expression, comprising any of the 27,400 fully defined nucleotide  
 sequences in the specification, or their complements or fragments, and  
 encoding at least 8 amino acids of any of the 688 amino acid sequences  
 fully defined in the specification. The probe is a single exon probe that  
 hybridizes under high stringency conditions to a nucleic acid molecule  
 expressed in human cells or tissues. Also included are a spatially-  
 addressable set of single exon nucleic acid probes for measuring human  
 gene expression (comprising a plurality of single exon nucleic acid  
 probes cited above, where each of the plurality of probes is separately  
 and addressably isolatable or amplifiable from the plurality), a single  
 exon microarray for measuring human gene expression, a method of  
 measuring human gene expression, a vector comprising the single exon  
 probe cited above, an ORF-encoded peptide comprising at least 8  
 contiguous amino acids of any of the above-mentioned amino acid  
 sequences (optionally with conservative amino acid substitutions), an  
 isolated antibody that binds specifically to a peptide cited above,  
 a method of selling and/or licensing single exon probes or microarrays to  
 a customer desiring to measure gene expression, a method of providing  
 human gene expression data by subscription, and a computer-readable  
 storage medium which contains a database having a plurality of records  
 (each record including data on the expression of a single exon probe  
 cited above). The probe, methods and apparatus are useful in gene  
 expression analysis. The probes may be used as tools for surveying  
 tissues to detect the presence of expressed messages that contain their  
 specific exon, or in constructing genome-derived single exon microarrays.  
 In addition, the probes are used in identifying and characterizing  
 alternative splicing events, in detecting and characterizing gross  
 alterations in the genomic locus that includes their exon, in assessing

CC smaller genomic alterations, in priming the synthesis of nucleic acids,  
 or in expressing the ORF-encoded peptide. The present sequence is a human  
 single exon probe protein of the invention. Note: The sequence data for  
 this patent did not form part of the printed specification, but was  
 obtained in electronic format directly from USPTO at  
 seqdata.uspto.gov/sequence.html?DocID=20030194704

XX  
 SQ Sequence 32 AA;  
 Query Match 29.9%; Score 29; DB 8; Length 32;  
 Best Local Similarity 71.4%; Pred. No. 1.3e+03;  
 Matches 5; Conservative 2; Mismatches 0; Indels 0; Gaps 0;

QY 12 KIVSQEP 18  
 Db 17 KIVSQEP 23  
 : |||||  
 : |||||

RESULT 142  
 AAM42284  
 ID AAM42284 standard; protein; 33 AA.  
 XX AC AAM42284;  
 XX DT 19-OCT-2001 (first entry)  
 XX DE Human breast or ovarian antigen SEQ ID NO: 161.  
 XX Human; breast antigen; ovarian antigen; cancer; metastasis; gene therapy.  
 XX OS Homo sapiens.  
 XX PN WO200155324-A2.  
 XX PD 02-AUG-2001.  
 XX PF 17-JAN-2001; 2001WO-US001344.  
 XX PR 31-JAN-2000; 2000US-0179065P.  
 PR 04-FEB-2000; 2000US-0180628P.  
 PR 24-FEB-2000; 2000US-0184664P.  
 PR 02-MAR-2000; 2000US-0186350P.  
 PR 16-MAR-2000; 2000US-0189874P.  
 PR 17-MAR-2000; 2000US-0190076P.  
 PR 18-APR-2000; 2000US-0198123P.  
 PR 19-MAY-2000; 2000US-0205515P.  
 PR 07-JUN-2000; 2000US-0209467P.  
 PR 28-JUN-2000; 2000US-0214886P.  
 PR 30-JUN-2000; 2000US-0215135P.  
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 PR 26-JUL-2000; 2000US-0220963P.  
 PR 26-JUL-2000; 2000US-0220964P.  
 PR 14-AUG-2000; 2000US-0224518P.  
 PR 14-AUG-2000; 2000US-0224519P.  
 PR 14-AUG-2000; 2000US-0225213P.  
 PR 14-AUG-2000; 2000US-0225214P.  
 PR 14-AUG-2000; 2000US-0225266P.  
 PR 14-AUG-2000; 2000US-0225267P.  
 PR 14-AUG-2000; 2000US-0225268P.  
 PR 14-AUG-2000; 2000US-0225270P.  
 PR 14-AUG-2000; 2000US-0225447P.  
 PR 14-AUG-2000; 2000US-0225757P.  
 PR 14-AUG-2000; 2000US-0225758P.  
 PR 14-AUG-2000; 2000US-0225759P.  
 PR 18-AUG-2000; 2000US-0226279P.  
 PR 22-AUG-2000; 2000US-0226681P.  
 PR 22-AUG-2000; 2000US-0226686P.  
 PR 23-AUG-2000; 2000US-0227182P.  
 PR 23-AUG-2000; 2000US-0227009P.

PR 30-AUG-2000; 2000US-0228924P.  
PR 01-SEP-2000; 2000US-0229287P.  
PR 01-SEP-2000; 2000US-0229343P.  
PR 01-SEP-2000; 2000US-0229344P.  
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PR 05-SEP-2000; 2000US-0229509P.  
PR 05-SEP-2000; 2000US-0229513P.  
PR 06-SEP-2000; 2000US-0230437P.  
PR 06-SEP-2000; 2000US-0230438P.  
PR 08-SEP-2000; 2000US-0231242P.  
PR 08-SEP-2000; 2000US-0231243P.  
PR 08-SEP-2000; 2000US-0231244P.  
PR 08-SEP-2000; 2000US-0231413P.  
PR 08-SEP-2000; 2000US-0231414P.  
PR 08-SEP-2000; 2000US-0232080P.  
PR 08-SEP-2000; 2000US-0232081P.  
PR 12-SEP-2000; 2000US-0231968P.  
PR 14-SEP-2000; 2000US-0232397P.  
PR 14-SEP-2000; 2000US-0232398P.  
PR 14-SEP-2000; 2000US-0232399P.  
PR 14-SEP-2000; 2000US-0232400P.  
PR 14-SEP-2000; 2000US-0232401P.  
PR 14-SEP-2000; 2000US-0233063P.  
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PR 21-SEP-2000; 2000US-0234223P.  
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PR 25-SEP-2000; 2000US-0234997P.  
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PR 29-SEP-2000; 2000US-0236367P.  
PR 29-SEP-2000; 2000US-0236368P.  
PR 29-SEP-2000; 2000US-0236369P.  
PR 29-SEP-2000; 2000US-0236370P.  
PR 02-OCT-2000; 2000US-0236802P.  
PR 02-OCT-2000; 2000US-0237037P.  
PR 02-OCT-2000; 2000US-0237038P.  
PR 02-OCT-2000; 2000US-0237039P.  
PR 02-OCT-2000; 2000US-0237040P.  
PR 13-OCT-2000; 2000US-0239935P.  
PR 13-OCT-2000; 2000US-0239937P.  
PR 20-OCT-2000; 2000US-0240960P.  
PR 20-OCT-2000; 2000US-0241221P.  
PR 20-OCT-2000; 2000US-0241785P.  
PR 20-OCT-2000; 2000US-0241786P.  
PR 20-OCT-2000; 2000US-0241787P.  
PR 20-OCT-2000; 2000US-0241808P.  
PR 20-OCT-2000; 2000US-0241809P.  
PR 20-OCT-2000; 2000US-0241826P.  
PR 01-NOV-2000; 2000US-0244617P.  
PR 08-NOV-2000; 2000US-0246474P.  
PR 08-NOV-2000; 2000US-0246475P.  
PR 08-NOV-2000; 2000US-0246476P.  
PR 08-NOV-2000; 2000US-0246477P.  
PR 08-NOV-2000; 2000US-0246478P.  
PR 08-NOV-2000; 2000US-0246523P.  
PR 08-NOV-2000; 2000US-0246524P.  
PR 08-NOV-2000; 2000US-0246525P.  
PR 08-NOV-2000; 2000US-0246526P.  
PR 08-NOV-2000; 2000US-0246527P.  
PR 08-NOV-2000; 2000US-0246528P.  
PR 08-NOV-2000; 2000US-0246532P.  
PR 08-NOV-2000; 2000US-0246609P.  
PR 08-NOV-2000; 2000US-0246610P.  
PR 08-NOV-2000; 2000US-0246611P.  
PR 08-NOV-2000; 2000US-0246613P.  
PR 17-NOV-2000; 2000US-0249207P.  
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PR 17-NOV-2000; 2000US-0249213P.  
PR 17-NOV-2000; 2000US-0249214P.  
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PR 17-NOV-2000; 2000US-0249216P.  
PR 17-NOV-2000; 2000US-0249217P.  
PR 17-NOV-2000; 2000US-0249218P.  
PR 17-NOV-2000; 2000US-0249244P.  
PR 17-NOV-2000; 2000US-0249245P.  
PR 17-NOV-2000; 2000US-0249264P.  
PR 17-NOV-2000; 2000US-0249265P.  
PR 17-NOV-2000; 2000US-0249297P.  
PR 17-NOV-2000; 2000US-0249299P.  
PR 17-NOV-2000; 2000US-0249300P.  
PR 01-DEC-2000; 2000US-0250160P.  
PR 01-DEC-2000; 2000US-0250391P.  
PR 05-DEC-2000; 2000US-0251030P.  
PR 05-DEC-2000; 2000US-0251988P.  
PR 05-DEC-2000; 2000US-0256719P.  
PR 06-DEC-2000; 2000US-0251479P.  
PR 08-DEC-2000; 2000US-0251856P.  
PR 08-DEC-2000; 2000US-0251868P.  
PR 08-DEC-2000; 2000US-0251869P.  
PR 08-DEC-2000; 2000US-0251989P.  
PR 08-DEC-2000; 2000US-0251990P.  
PR 11-DEC-2000; 2000US-0254097P.  
PR 05-JAN-2001; 2001US-0259678P.  
XX  
FA (HUMA-) HUMAN GENOME SCI INC.  
XX  
XX Rosen CA, Barash SC, Ruben SM;  
XX  
XX WPI; 2001-488785/53.  
DR N-PSDB; AAI62511.  
XX  
PT New isolated nucleic acids and polypeptides, useful for diagnosing,  
PT treating and/or preventing human diseases and disorders.  
XX  
XX Claim 11; SEQ ID NO 161; 520pp + Sequence Listing; English.  
PS  
XX The present invention provides the protein and coding sequences of a  
CC number of ovarian and breast antigens. These are shown in AAI62467-  
CC AAI62572 and AAM42240-AA42345. The sequences can be used in the  
CC diagnosis, prevention and treatment of breast and ovarian cancers, and  
CC their metastases. The present sequence is a protein of the invention.  
CC Note: The sequence data for this patent did not form part of the printed  
CC specification, but was obtained in electronic format directly from WIPO  
CC at ftp.wipo.int/pub/published\_pct\_sequences  
XX  
SQ Sequence 33 AA;  
Query March 29.9%; Score 29; DB 4; Length 33;  
Best Local Similarity 50.0%; Pred.No. 1.4e+03;  
Matches 5; Conservative 2; Mismatches 3; Indels 0; Gaps 0;  
Oy 2 PNHLSKIAF 11  
Db 14 PNKLTSQUTF 23  
RESULT 143  
AAM96243  
ID AAM96243 standard; protein; 33 AA.  
XX AAM96243;  
XX  
XX 21-NOV-2001 (first entry)  
XX Human reproductive system related antigen SEQ ID NO: 4901.  
XX Human; reproductive system related antigen; reproductive system disorder;  
KW cancer; gene therapy.



OS  
XX Homo sapiens.  
PN WO200155320-A2.  
XX  
XX  
XX 02-AUG-2001.  
XX  
XX 17-JAN-2001; 2001WO-US001339.  
XX  
XX 31-JAN-2000; 2000US-0179065P.  
XX 04-FEB-2000; 2000US-0180628P.  
XX 24-FEB-2000; 2000US-0184664P.  
XX 02-MAR-2000; 2000US-0186350P.  
XX 16-MAR-2000; 2000US-0189874P.  
XX 17-MAR-2000; 2000US-0190076P.  
XX 18-APR-2000; 2000US-0198123P.  
XX 19-MAY-2000; 2000US-0205515P.  
XX 07-JUN-2000; 2000US-0209467P.  
XX 28-JUN-2000; 2000US-0214886P.  
XX 30-JUN-2000; 2000US-0215135P.  
XX 07-JUL-2000; 2000US-0216647P.  
XX 07-JUL-2000; 2000US-0216880P.  
XX 11-JUL-2000; 2000US-0217487P.  
XX 11-JUL-2000; 2000US-0217496P.  
XX 14-JUL-2000; 2000US-0218290P.  
XX 26-JUL-2000; 2000US-0220963P.  
XX 26-JUL-2000; 2000US-0220964P.  
XX 14-AUG-2000; 2000US-0224518P.  
XX 14-AUG-2000; 2000US-0224519P.  
XX 14-AUG-2000; 2000US-0225213P.  
XX 14-AUG-2000; 2000US-0225214P.  
XX 14-AUG-2000; 2000US-0225266P.  
XX 14-AUG-2000; 2000US-0225267P.  
XX 14-AUG-2000; 2000US-0225268P.  
XX 14-AUG-2000; 2000US-0225270P.  
XX 14-AUG-2000; 2000US-0225447P.  
XX 14-AUG-2000; 2000US-0225757P.  
XX 14-AUG-2000; 2000US-0225758P.  
XX 14-AUG-2000; 2000US-0225759P.  
XX 18-AUG-2000; 2000US-0226279P.  
XX 22-AUG-2000; 2000US-0226681P.  
XX 22-AUG-2000; 2000US-0226868P.  
XX 22-AUG-2000; 2000US-0227182P.  
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XX 30-AUG-2000; 2000US-0228924P.  
XX 01-SEP-2000; 2000US-0229287P.  
XX 01-SEP-2000; 2000US-0229343P.  
XX 01-SEP-2000; 2000US-0229344P.  
XX 01-SEP-2000; 2000US-0229345P.  
XX 05-SEP-2000; 2000US-0229509P.  
XX 05-SEP-2000; 2000US-0229513P.  
XX 06-SEP-2000; 2000US-0230437P.  
XX 06-SEP-2000; 2000US-0230438P.  
XX 08-SEP-2000; 2000US-0231242P.  
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XX 08-SEP-2000; 2000US-0231413P.  
XX 08-SEP-2000; 2000US-0231414P.  
XX 08-SEP-2000; 2000US-0232080P.  
XX 08-SEP-2000; 2000US-0232081P.  
XX 12-SEP-2000; 2000US-0231968P.  
XX 14-SEP-2000; 2000US-0232397P.  
XX 14-SEP-2000; 2000US-0232398P.  
XX 14-SEP-2000; 2000US-0232399P.  
XX 14-SEP-2000; 2000US-0232400P.  
XX 14-SEP-2000; 2000US-0232401P.  
XX 14-SEP-2000; 2000US-0233063P.  
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XX 21-SEP-2000; 2000US-0233065P.  
XX 21-SEP-2000; 2000US-0234223P.  
XX 25-SEP-2000; 2000US-0234274P.  
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PR 26-SEP-2000; 2000US-0235484P.  
PR 27-SEP-2000; 2000US-0235834P.  
PR 27-SEP-2000; 2000US-0235836P.  
PR 29-SEP-2000; 2000US-0236327P.  
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PR 29-SEP-2000; 2000US-0236368P.  
PR 29-SEP-2000; 2000US-0236369P.  
PR 29-SEP-2000; 2000US-0236370P.  
PR 02-OCT-2000; 2000US-0236802P.  
PR 02-OCT-2000; 2000US-0237037P.  
PR 02-OCT-2000; 2000US-0237038P.  
PR 02-OCT-2000; 2000US-0237039P.  
PR 02-OCT-2000; 2000US-0237040P.  
PR 13-OCT-2000; 2000US-0239935P.  
PR 13-OCT-2000; 2000US-0239937P.  
PR 20-OCT-2000; 2000US-0240960P.  
PR 20-OCT-2000; 2000US-0241221P.  
PR 20-OCT-2000; 2000US-0241785P.  
PR 20-OCT-2000; 2000US-0241786P.  
PR 20-OCT-2000; 2000US-0241787P.  
PR 20-OCT-2000; 2000US-0241808P.  
PR 20-OCT-2000; 2000US-0241809P.  
PR 20-OCT-2000; 2000US-0241826P.  
PR 01-NOV-2000; 2000US-0244617P.  
PR 08-NOV-2000; 2000US-0246474P.  
PR 08-NOV-2000; 2000US-0246475P.  
PR 08-NOV-2000; 2000US-0246476P.  
PR 08-NOV-2000; 2000US-0246477P.  
PR 08-NOV-2000; 2000US-0246478P.  
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PR 08-NOV-2000; 2000US-0246524P.  
PR 08-NOV-2000; 2000US-0246525P.  
PR 08-NOV-2000; 2000US-0246526P.  
PR 08-NOV-2000; 2000US-0246527P.  
PR 08-NOV-2000; 2000US-0246528P.  
PR 08-NOV-2000; 2000US-0246532P.  
PR 08-NOV-2000; 2000US-0246609P.  
PR 08-NOV-2000; 2000US-0246610P.  
PR 08-NOV-2000; 2000US-0246611P.  
PR 08-NOV-2000; 2000US-0246613P.  
PR 17-NOV-2000; 2000US-0249207P.  
PR 17-NOV-2000; 2000US-0249208P.  
PR 17-NOV-2000; 2000US-0249209P.  
PR 17-NOV-2000; 2000US-0249210P.  
PR 17-NOV-2000; 2000US-0249211P.  
PR 17-NOV-2000; 2000US-0249212P.  
PR 17-NOV-2000; 2000US-0249213P.  
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PR 17-NOV-2000; 2000US-0249217P.  
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PR 17-NOV-2000; 2000US-0249244P.  
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PR 17-NOV-2000; 2000US-0249264P.  
PR 17-NOV-2000; 2000US-0249265P.  
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PR 17-NOV-2000; 2000US-0249299P.  
PR 17-NOV-2000; 2000US-0249300P.  
PR 01-DEC-2000; 2000US-0250160P.  
PR 01-DEC-2000; 2000US-0250391P.  
PR 05-DEC-2000; 2000US-0251030P.  
PR 05-DEC-2000; 2000US-0251988P.  
PR 05-DEC-2000; 2000US-0256719P.  
PR 06-DEC-2000; 2000US-0251479P.  
PR 08-DEC-2000; 2000US-0251856P.  
PR 08-DEC-2000; 2000US-0251868P.  
PR 08-DEC-2000; 2000US-0251869P.  
PR 08-DEC-2000; 2000US-0251989P.  
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PR 11-DEC-2000; 2000US-0254097P.  
PR 05-JAN-2001; 2001US-0259678P.  
XX

PA (HUMA-) HUMAN GENOME SCI INC.  
XX  
PI Rosen CA, Barash SC, Ruben SM;  
XX  
DR WPI; 2001-465570/50.  
DR N-PSDB; AAL02213.  
XX  
PT Isolated nucleic acid molecule encoding a reproductive system antigen is  
PT used in preventing, treating or ameliorating a medical condition.  
XX  
PS Claim 11; SEQ ID NO 4901; 1297pp + Sequence Listing; English.  
XX  
CC The present invention provides the protein and coding sequences of a  
CC number of human reproductive system related antigens. These can be used  
CC in the prevention and treatment of reproductive system disorders,  
CC including cancer. The present sequence is a protein of the invention  
XX  
XX Sequence 33 AA;  
XX  
XX Query Match 29.9%; Score 29; DB 4; Length 33;  
XX Best Local Similarity 50.0%; Pred. No. 1.4e+03;  
XX Matches 5; Conservative 2; Mismatches 3; Indels 0; Gaps 0;  
XX  
QY 2 PNHLNSKIAP 11  
DB ||| ||:|  
14 PNKLTSQTLF 23  
XX  
RESULT 144  
ADE01861  
ID ADE01861 standard; peptide; 33 AA.  
XX  
AC ADE01861;  
XX  
DT 29-JAN-2004 (first entry)  
XX  
DE Hybrid polypeptide pharmacokinetic enhancer peptide, SEQ ID No 368.  
XX  
XX hybrid; enhancer; anti-fusogenic; antiviral; virucide; antidiabetic;  
XX pharmacokinetic; fusogenic; insulin; diabetes.  
XX  
OS Unidentified.  
XX  
FH Key Location/Qualifiers  
FT Modified-site 1  
FT /note= "Residue is modified by acetyl group"  
FT Modified-site 33  
FT /note= "C-terminal amide"  
XX  
PN US6348568-B1.  
XX  
XX 19-FEB-2002.  
XX  
XX 20-MAY-1999; 99US-00315304.  
PF  
XX 20-MAY-1998; 98US-00082279.  
PR  
XX (TRIM-) TRIMERIS INC.  
XX  
XX Barney S, Guthrie KI, Merutka G, Anwer MK, Lambert DM;  
XX  
XX WPI; 2002-424396/45.  
XX  
XX New hybrid polypeptide for modulating fusogenic events for e.g. antiviral  
PT activity, has enhancer peptide sequence derived from retroviral envelope  
PT protein sequences linked to core polypeptide e.g. therapeutic protein.  
PT  
XX Disclosure; SEQ ID NO 368; 70pp; English.  
XX  
XX The invention relates to a novel hybrid polypeptide comprising an  
CC enhancer peptide sequence linked to a core polypeptide. The enhancer  
CC peptide sequence comprises QWEQEKI or WASLWEP. The invention also  
CC includes novel peptides that exhibit anti-fusogenic activity, antiviral

CC dedifferentiation comprising inhibiting the interaction of Nef with a Src  
CC family tyrosine kinase SH3 domain of a polypeptide of the cell. The  
CC method is useful for inhibiting kidney cell dedifferentiation for  
CC treating e.g., HIV associated nephropathy (HIVAN), AIDS, dementia,  
CC anaemia, lymphoma, myopathy, cardiomyopathy or primary HIV-induced  
CC disease progression. The present sequence represents the amino acid  
CC sequence of a Nef/SH3 inhibitory peptide.

Query Match	29.9%	Score 29;	DB 8;	Length 33;
Best Local Similarity	41.7%	Pred. No.	1.4e+03;	
Matches 5;	Conservative	2;	Mismatches 5;	Indels 0;
Gaps	0;			

Qy           8 KIAFKIVSQEPA 19  
               | : | : |  
Db          17 KVGFPVTPOVPA 28

RESULT 146	
AAW30694	
ID	AAW30694 standard; peptide; 34 AA.
XX	
XX	
AC	AAW30694;
XX	
XX	
DT	11-JAN-1999 (first entry)
XX	
XX	
DE	Repro-PC-1.0 immunogenic peptide 42.
XX	
KW	Repro-PC-1.0; prostate cancer; marker; synaptotagmin; human; diagnosis;
KW	vaccine; therapy; immunogen.
XX	
OS	Homo sapiens.

Query Match	29.9%;	Score 29;	DB 2;	Length 34;
Best Local Similarity	60.0%;	Pred. No. 1.4e+03;		
Matches	6;	Conservative	1;	Mismatches 3; Indels 0; Gaps 0;
Qy	2	PNHLNSKIAF	11	
		:		
Db	23	PENLNSKKKF	32	

Qy	2	PNHLSKIAF	11
		:	
Db	23	PENLSKKKF	32

RESULT 147	
AAy89007	
XX	AAy89007 standard; peptide; 34 AA.
XX	
AC	AAy89007;
XX	
DT	23-MAY-2000 (first entry)
XX	
DE	Core polypeptide fragment T No. 371.
XX	
KW	Retrovirus; hybrid polypeptide; enhancer; gp41; envelope protein; HIV-1;
KW	HIV-2; SV; pharmacokinetic; half-life; growth factor; cytokine; viral;
KW	anti-fusogenic; differentiation factor; interleukin; interferon;
KW	colony stimulating factor; hormone; angiogenic factor.
XX	
OS	Unidentified.

Query Match	29.9%	Score 29;	DB 3;	Length 34;
Best Local Similarity	40.0%	Pred. No.	1.4e+03;	
Matches	6;	Conservative	2;	Mismatches 7; Indels 0; Gaps 0;

Sequence 34 AA;

QY 3 NHLNSKIAFKIVSQE 17  
 Db 15 NKXNGTDAVKLIKQE 29

RESULT 148  
 AAY89008  
 ID AAY89008 standard; peptide; 34 AA.  
 XX  
 AC AAY89008;  
 XX

DT 23-MAY-2000 (first entry)  
 DE Core polypeptide fragment T No. 372.  
 XX

KW Retrovirus; hybrid polypeptide; enhancer; gp41; envelope protein; HIV-1;  
 KW HIV-2; SIV; pharmacokinetic; half-life; growth factor; cytokine; viral;  
 KW anti-fusogenic; differentiation factor; interleukin; interferon;  
 KW colony stimulating factor; hormone; angiogenic factor.  
 XX

OS Unidentified.  
 XX  
 XX  
 FH Key Location/Qualifiers  
 FT Misc-difference 16 /note= "indicated as U in the specification"  
 FT  
 XX

PN WO9959615-A1.

XX 25-NOV-1999.

XX 20-MAY-1999; 99WO-US011219.

XX 20-MAY-1998; 98US-00082279.

XX (TRIM-) TRIMERIS INC.

XX Barney S, Guthrie KI, Merutka G, Anwer MK, Lambert DM;  
 PI WPI; 2000-136792/12.

XX A new hybrid polypeptide with enhanced pharmacokinetic properties  
 PT comprises enhancer sequence.  
 PT

XX Disclosure; Page 28; 124pp; English.

XX The invention relates to hybrid polypeptides comprising enhancer peptide  
 CC sequence linked to core polypeptides. The enhancer polypeptides are  
 CC derived from various retroviral envelope (gp41) protein sequences,  
 CC especially from HIV-1, HIV-2 and SIV. The enhancer peptides enhance the  
 CC pharmacokinetic properties such as increasing the half-life of any core  
 CC polypeptide that they are linked to. The core polypeptides are any  
 CC polypeptide that may be introduced into a living system and that can  
 CC function as a pharmacologically useful peptide for the treatment or  
 CC prevention of a disease. The core polypeptides are bioactive peptides  
 CC selected from a growth factor, cytokine, differentiation factor,  
 CC interleukin, interferon, colony stimulating factor, hormone or angiogenic  
 CC factor. The peptides of the invention can be used for inhibiting viral  
 CC infection and can be used in anti-viral and anti-fusogenic treatments.  
 CC Sequences AAY8651-Y90055 represent core polypeptide fragments that can  
 CC be used in the invention. Some sequences among those indicated also  
 CC comprise enhancer fragments at terminal ends and form hybrid polypeptides  
 XX

XX Sequence 34 AA;

Query Match 29.9%; Score 29; DB 3; Length 34;  
 Best Local Similarity 40.0%; Pred. No. 1.4e+03;  
 Matches 6; Conservative 2; Mismatches 7; Indels 0; Gaps 0;

QY 3 NHLNSKIAFKIVSQE 17  
 Db 14 NKXNGTDAVKLIKQE 28

RESULT 149

AAY89009  
 ID AAY89009 standard; peptide; 34 AA.  
 XX

AC AAY89009;

XX 23-MAY-2000 (first entry)

XX Core polypeptide fragment T No. 373.

KW Retrovirus; hybrid polypeptide; enhancer; gp41; envelope protein; HIV-1;  
 KW HIV-2; SIV; pharmacokinetic; half-life; growth factor; cytokine; viral;  
 KW anti-fusogenic; differentiation factor; interleukin; interferon;  
 KW colony stimulating factor; hormone; angiogenic factor.  
 XX

OS Unidentified.

XX Key Location/Qualifiers

FT Misc-difference 15 /note= "indicated as U in the specification"

XX WO9959615-A1.

XX 25-NOV-1999.

XX 20-MAY-1999; 99WO-US011219.

XX 20-MAY-1998; 98US-00082279.

XX (TRIM-) TRIMERIS INC.

XX Barney S, Guthrie KI, Merutka G, Anwer MK, Lambert DM;  
 PI WPI; 2000-136792/12.

XX A new hybrid polypeptide with enhanced pharmacokinetic properties  
 PT comprises enhancer sequence.  
 PT

XX Disclosure; Page 28; 124pp; English.

XX The invention relates to hybrid polypeptides comprising enhancer peptide  
 CC sequence linked to core polypeptides. The enhancer polypeptides are  
 CC derived from various retroviral envelope (gp41) protein sequences,  
 CC especially from HIV-1, HIV-2 and SIV. The enhancer peptides enhance the  
 CC pharmacokinetic properties such as increasing the half-life of any core  
 CC polypeptide that they are linked to. The core polypeptides are any  
 CC polypeptide that may be introduced into a living system and that can  
 CC function as a pharmacologically useful peptide for the treatment or  
 CC prevention of a disease. The core polypeptides are bioactive peptides  
 CC selected from a growth factor, cytokine, differentiation factor,  
 CC interleukin, interferon, colony stimulating factor, hormone or angiogenic  
 CC factor. The peptides of the invention can be used for inhibiting viral  
 CC infection and can be used in anti-viral and anti-fusogenic treatments.  
 CC Sequences AAY8651-Y90055 represent core polypeptide fragments that can  
 CC be used in the invention. Some sequences among those indicated also  
 CC comprise enhancer fragments at terminal ends and form hybrid polypeptides  
 XX

XX Sequence 34 AA;

Query Match 29.9%; Score 29; DB 3; Length 34;  
 Best Local Similarity 40.0%; Pred. No. 1.4e+03;  
 Matches 6; Conservative 2; Mismatches 7; Indels 0; Gaps 0;

QY 3 NHLNSKIAFKIVSQE 17

Db 13 NKXNGTDAVKLIKQE 27

RESULT 150

AAB54800  
 ID AAB54800 standard; peptide; 34 AA.  
 XX

XX AAB54800;

XX 05-MAR-2001 (first entry)  
 DT RSV antiviral activity exhibiting peptide SEQ ID NO:17.  
 DE  
 XX Long lasting fusion peptide inhibitor; viral infection; antiviral;  
 XX antifusogenic; mobile blood component; measles virus; MeV; SIV;  
 KW smian immunodeficiency virus; human parainfluenza virus; HPiV; RSV;  
 KW human respiratory syncytial virus; human immunodeficiency virus; HIV.  
 XX  
 OS Human respiratory syncytial virus.  
 XX  
 XX WO200069902-A1.  
 XX  
 XX 23-NOV-2000.  
 XX  
 XX 17-MAY-2000; 2000WO-US013651.  
 PF  
 XX 17-MAY-1999; 99US-0134406P.  
 PR  
 XX 10-SEP-1999; 99US-0153406P.  
 PR  
 XX (CONJ-) CONJUCHEM INC.  
 PA  
 XX Bridon DP, Dufresne RP, Boudjellab N, Robitaille M, Milner PG;  
 PI WPI; 2001-007496/01.  
 XX  
 XX A modified peptide and a reactive group which is reactive with amino  
 PT groups, hydroxyl groups, or thiol groups on blood components to form  
 PT stable covalent bonds useful for treatment of viral infections, e.g.  
 PT human immunodeficiency virus.  
 PT  
 PS Claim 9; Page 179; 21pp; English.  
 XX  
 CC The present invention describes a modified anti-viral peptide (I)  
 CC comprising a peptide that exhibits anti-viral activity and a reactive  
 CC group which is reactive with amino groups, hydroxyl groups, or thiol  
 CC groups on blood components to form stable covalent bonds. (I) has anti-  
 CC viral and anti-fusogenic activities. (I) inhibits viral infection of  
 CC cells by inhibiting cell-cell fusion or free virus infection or to reduce  
 CC the level of membrane fusion events between two or more entities, e.g.,  
 CC virus-cell or cell-cell, relative to the level of membrane fusion that  
 CC occurs in the absence of the peptide. (I) is useful in the treatment of  
 CC patients who are suffering from viral infection, e.g. HIV, RSV, HPiV,  
 CC MeV, and SIV. (I) may be administered prophylactically to previously,  
 CC uninfected individuals. This is useful in cases where an individual has  
 CC been subjected to a high risk of exposure to a virus. By bonding of long-  
 CC lived components of the blood, such as immunoglobulin, serum albumin, red  
 CC blood cells and platelets the activity is extended for days to weeks.  
 CC This is due to improved stability in vivo and a reduced susceptibility to  
 CC peptidase or protease degradation. This minimises the need for more  
 CC frequent, or even continual, administration of the peptides. AAB54784 to  
 CC AAB55431 represent peptides used in the exemplification of the present  
 CC invention  
 XX  
 SQ Sequence 34 AA;  
 Query Match 29.9%; Score 29; DB 4; Length 34;  
 Best Local Similarity 33.3%; Pred. No. 1.4e+03;  
 Matches 5; Conservative 3; Mismatches 7; Indels 0; Gaps 0;  
 QY 3 NHLNSKIAPKIVSQE 17  
 Db 12 NKNMGDAKVKLIKQE 26  
 RESULT 151  
 AAB92259  
 ID AAB92259 standard; peptide; 34 AA.  
 XX  
 AC AAB92259;  
 XX  
 DT 22-JUN-2001 (first entry)

XX Virus related peptide SEQ ID NO:1435.  
 DE  
 XX Protection; endogenous therapeutic peptide; peptidase; conjugation;  
 KW blood component; modification; succinimidyl; maleimido group; amino;  
 KW hydroxyl; thiol; hormone; growth factor; neurotransmitter.  
 XX  
 OS Homo sapiens.  
 OS Synthetic.  
 XX  
 PN WO200069900-A2.  
 XX  
 XX 23-NOV-2000.  
 XX  
 XX 17-MAY-2000; 2000WO-US013576.  
 PF  
 XX 17-MAY-1999; 99US-0134406P.  
 PR  
 XX 10-SEP-1999; 99US-0153406P.  
 PR  
 XX 15-OCT-1999; 99US-0159783P.  
 XX  
 PA (CONJ-) CONJUCHEM INC.  
 XX  
 XX Bridon DP, Ezrin AM, Milner PG, Holmes DL, Thibaudeau K;  
 PI WPI; 2001-112059/12.  
 XX  
 XX Modifying and attaching therapeutic peptides to albumin prevents  
 PT peptidase degradation, useful for increasing length of in vivo activity.  
 PT  
 PS Disclosure; Page 668; 733pp; English.  
 XX  
 CC The present invention describes a modified therapeutic peptide (I)  
 CC comprising a therapeutically active amino acid region (III) and a  
 CC reactive group (II) (e.g. succinimidyl and maleimido groups) attached to  
 CC a less therapeutically active amino acid region (IV), which covalently  
 CC bonds with amino/hydroxyl/thiol groups on blood components to form a  
 CC peptidase stabilised therapeutic peptide composed of 3-50 amino acids.  
 CC (I) are useful for modifying therapeutic peptides e.g. hormones, growth  
 CC factors and neurotransmitters, to protect them from peptidase activity in  
 CC vivo for the treatment of various disorders. Endogenous therapeutic  
 CC peptides are not suitable as drug candidates as they require frequent  
 CC administration due to rapid degradation by peptidases in the body.  
 CC Modifying and attaching therapeutic peptides to albumin prevents or  
 CC reduces the action of peptidases to increase length of activity (half  
 CC life) and specificity as bonding to large molecules decreases  
 CC intracellular uptake and interference with physiological processes.  
 CC AAB90829 to AAB92441 represent peptides which can be used in the  
 CC exemplification of the present invention  
 XX  
 SQ Sequence 34 AA;  
 Query Match 29.9%; Score 29; DB 4; Length 34;  
 Best Local Similarity 33.3%; Pred. No. 1.4e+03;  
 Matches 5; Conservative 3; Mismatches 7; Indels 0; Gaps 0;  
 QY 3 NHLNSKIAPKIVSQE 17  
 Db 12 NKNMGDAKVKLIKQE 26  
 RESULT 152  
 AAB77362  
 ID AAB77362 standard; peptide; 34 AA.  
 XX  
 AC AAB77362;  
 XX  
 DT 19-APR-2001 (first entry)  
 XX  
 DE Core polypeptide T371.  
 XX  
 KW Core polypeptide; enhancer; antiviral; anti-HIV; virucide; hepatotropic;  
 KW antiinflammatory; hybrid polypeptide; coiled-coil peptide interaction;  
 KW fusion-related disorder; bacterial infection; viral infection.

```

XX OS Unidentified.
XX PN WO200103723-A1.
XX PD 18-JAN-2001.
XX PF 10-JUL-2000; 2000WO-US018772.
XX PR 09-JUL-1999; 99US-00350641.
XX PA (TRIM-) TRIMERIS INC.
XX PI Barney S, Guthrie KI, Merutka G, Anwer MK, Lambert DM;
XX WPI; 2001-147136/15.
XX
XX New hybrid polypeptide, useful for preventing, treating and diagnosing
XX e.g. viral infections, comprises an enhancer peptide linked to a core
XX polypeptide.
XX
XX Disclosure; Page 38; 15lpp; English.
XX
XX The present sequence is a core polypeptide which may be linked to an
XX enhancer peptide to form a novel hybrid polypeptide. The hybrid
XX polypeptide exhibits enhanced pharmacokinetic properties relative to
XX those exhibited by the core polypeptide when introduced into a living
XX system. It is used to increase the in vitro or ex vivo half-life of the
XX core polypeptide. The hybrid and core polypeptides can be used for
XX modulating fucogenic events and intracellular processes involving coiled-
XX coil peptide interactions. Other uses include preventing, treating and/or
XX diagnosing disorders involving fusion events (e.g. modulation of
XX neurotransmitter exchange and sperm-egg fusion), intracellular processes
XX involving coiled-coil peptides (e.g. bacterial infections) and viral
XX infections that involve cell-cell and/or virus-cell fusion (e.g. viral
XX infections caused by human immunodeficiency virus, respiratory syncytial
XX virus, Epstein-Barr virus, hepatitis B virus, Mason-Pfizer virus and
XX polio virus). The enhancer peptide sequence increases the half-life and
XX reduces the clearance rate of therapeutic peptides, which increases their
XX efficacy and minimises the incidence and severity of adverse side
XX effects. In addition, this increases the sensitivity of the diagnostic
XX procedure in which they are used
XX
XX Sequence 34 AA;
XX
XX Query Match 29.9%; Score 29; DB 4; Length 34;
XX Best Local Similarity 40.0%; Pred. NO. 1.4e+03;
XX Matches 6; Conservative 2; Mismatches 7; Indels 0; Gaps 0;
XX
XX QY 3 NHLNSKIAFKIVSOE 17
XX DB 15 NXNGTDAVKLIKQE 29
XX
XX RESULT 153
XX AAB77364
XX ID AAB77364 standard; peptide; 34 AA.
XX AC AAB77364;
XX
XX 19-APR-2001 (first entry)
XX
XX Core polypeptide T373.
XX
XX Core polypeptide; enhancer; antiviral; anti-HIV; virucide; hepatotropic;
XX antiinflammatory; hybrid polypeptide; coiled-coil peptide interaction;
XX fusion-related disorder; bacterial infection; viral infection.
XX
XX Unidentified.
XX
XX WO200103723-A1.
XX
XX 18-JAN-2001.

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XX PF 10-JUL-2000; 2000WO-US018772.
XX PR 09-JUL-1999; 99US-00350641.
XX PA (TRIM-) TRIMERIS INC.
XX PI Barney S, Guthrie KI, Merutka G, Anwer MK, Lambert DM;
XX WPI; 2001-147136/15.
XX
XX New hybrid polypeptide, useful for preventing, treating and diagnosing
XX e.g. viral infections, comprises an enhancer peptide linked to a core
XX polypeptide.
XX
XX Disclosure; Page 38; 15lpp; English.
XX
XX The present sequence is a core polypeptide which may be linked to an
XX enhancer peptide to form a novel hybrid polypeptide. The hybrid
XX polypeptide exhibits enhanced pharmacokinetic properties relative to
XX those exhibited by the core polypeptide when introduced into a living
XX system. It is used to increase the in vitro or ex vivo half-life of the
XX core polypeptide. The hybrid and core polypeptides can be used for
XX modulating fucogenic events and intracellular processes involving coiled-
XX coil peptide interactions. Other uses include preventing, treating and/or
XX diagnosing disorders involving fusion events (e.g. modulation of
XX neurotransmitter exchange and sperm-egg fusion), intracellular processes
XX involving coiled-coil peptides (e.g. bacterial infections) and viral
XX infections that involve cell-cell and/or virus-cell fusion (e.g. viral
XX infections caused by human immunodeficiency virus, respiratory syncytial
XX virus, Epstein-Barr virus, hepatitis B virus, Mason-Pfizer virus and
XX polio virus). The enhancer peptide sequence increases the half-life and
XX reduces the clearance rate of therapeutic peptides, which increases their
XX efficacy and minimises the incidence and severity of adverse side
XX effects. In addition, this increases the sensitivity of the diagnostic
XX procedure in which they are used
XX
XX Sequence 34 AA;
XX
XX Query Match 29.9%; Score 29; DB 4; Length 34;
XX Best Local Similarity 40.0%; Pred. NO. 1.4e+03;
XX Matches 6; Conservative 2; Mismatches 7; Indels 0; Gaps 0;
XX
XX QY 3 NHLNSKIAFKIVSOE 17
XX DB 13 NXNGTDAVKLIKQE 27
XX
XX RESULT 154
XX AAB77363
XX ID AAB77363 standard; peptide; 34 AA.
XX AC AAB77363;
XX
XX 19-APR-2001 (first entry)
XX
XX Core polypeptide T372.
XX
XX Core polypeptide; enhancer; antiviral; anti-HIV; virucide; hepatotropic;
XX antiinflammatory; hybrid polypeptide; coiled-coil peptide interaction;
XX fusion-related disorder; bacterial infection; viral infection.
XX
XX Unidentified.
XX
XX WO200103723-A1.
XX
XX 18-JAN-2001.
XX
XX 10-JUL-2000; 2000WO-US018772.
XX
XX 09-JUL-1999; 99US-00350641.
XX
XX (TRIM-) TRIMERIS INC.

```

XX  
PI Barney S, Guthrie KI, Merutka G, Anwer MK, Lambert DM;  
XX WPI; 2001-147136/15.  
XX  
XX  
XX New hybrid polypeptide, useful for preventing, treating and diagnosing  
PT e.g. viral infections, comprises an enhancer peptide linked to a core  
PT polypeptide.  
XX  
XX Disclosure; Page 38; 151pp; English.  
XX  
XX The present sequence is a core polypeptide which may be linked to an  
CC enhancer peptide to form a novel hybrid polypeptide. The hybrid  
CC polypeptide exhibits enhanced pharmacokinetic properties relative to  
CC those exhibited by the core polypeptide when introduced into a living  
CC system. It is used to increase the in vitro or ex vivo half-life of the  
CC core polypeptide. The hybrid and core polypeptides can be used for  
CC modulating fusing events and intracellular processes involving coiled-  
CC coil peptide interactions. Other uses include preventing, treating and/or  
CC diagnosing disorders involving fusion events (e.g. modulation of  
CC neurotransmitter exchange and sperm-egg fusion), intracellular processes  
CC involving coiled-coil peptides (e.g. bacterial infections) and viral  
CC infections that involve cell-cell and/or virus-cell fusion (e.g. viral  
CC virus, Epstein-Barr virus, hepatitis B virus, Mason-Pfizer virus and  
CC polio virus). The enhancer peptide sequence increases the half-life and  
CC reduces the clearance rate of therapeutic peptides, which increases their  
CC efficacy and minimises the incidence and severity of adverse side  
CC effects. In addition, this increases the sensitivity of the diagnostic  
CC procedure in which they are used  
XX  
XX Sequence 34 AA;  
SQ

Query Match 29.9%; Score 29; DB 4; Length 34;  
Best Local Similarity 40.0%; Pred. No. 1.4e+03;  
Matches 6; Conservative 2; Mismatches 7; Indels 0; Gaps 0;

Qy 3 NHLNSKIAFKIVSQE 17  
| | | | |  
Db 14 NXNGTDAVKLIKQE 28

RESULT 155  
ABB00367  
ID ABB00367 standard; peptide; 34 AA.  
AC ABB00367;  
XX  
XX 03-JAN-2002 (first entry)  
XX  
XX RSV F2 protein DP178/107-like region peptide T372.  
XX  
XX Human immunodeficiency virus; HIV; respiratory syncytial virus; RSV;  
XX virucide; heptad repeat region; transmembrane protein; gp41; HR1; HR2;  
XX infection.  
XX  
XX Human respiratory syncytial virus.  
XX  
XX  
XX Key Location/Qualifiers  
FT Modified-site 1  
FT /note= "N-terminal is substituted by Ac"  
FT Modified-site 34  
FT /note= "C-terminal amide"  
XX  
XX WO200164013-A2.  
XX  
XX 07-SEP-2001.  
XX  
XX 07-FEB-2001; 2001WO-US003988.  
XX  
XX 29-FEB-2000; 2000US-00515965.  
XX  
XX (TRIM-) TRIMERIS INC.  
XX  
XX

XX  
PI Antczak JB, Delmedico MK, Erickson JB, Lambert DM, Sista P;  
XX WPI; 2001-514829/56.  
XX  
XX  
XX Heptad repeat region peptide analogs useful for inhibiting virus/cells  
PT fusion, useful for treating HIV and Respiratory Syncytial Virus  
PT infection.  
XX  
XX Example; Page 40; 587pp; English.  
XX  
XX The invention relates to isolated analogues of the heptad repeat region  
CC peptides DP178 and DP107. DP178 and DP107 correspond to amino acids 638-  
CC 673 (heptad repeat region HR2) and 558-595 (heptad repeat region HR1)  
CC respectively, of HIV-1LAI transmembrane protein gp41. The HR1 and HR2  
CC regions of proteins interact non-covalently with each other and/or with  
CC peptides derived from them. This interaction is required for normal  
CC infectivity of viruses such as RSV and HIV. The heptad repeat region  
CC peptide analogues may be used to inhibit respiratory syncytial virus  
CC (RSV) infection in a cell. They may also be used to inhibit HIV  
CC infection. The present sequence is a peptide provided in the  
CC specification  
XX  
XX Sequence 34 AA;  
SQ

Query Match 29.9%; Score 29; DB 4; Length 34;  
Best Local Similarity 40.0%; Pred. No. 1.4e+03;  
Matches 6; Conservative 2; Mismatches 7; Indels 0; Gaps 0;

Qy 3 NHLNSKIAFKIVSQE 17  
| | | | |  
Db 14 NXNGTDAVKLIKQE 28

RESULT 156  
ABB00368  
ID ABB00368 standard; peptide; 34 AA.  
AC ABB00368;  
XX  
XX 03-JAN-2002 (first entry)  
XX  
XX RSV F2 protein DP178/107-like region peptide T373.  
XX  
XX Human immunodeficiency virus; HIV; respiratory syncytial virus; RSV;  
XX virucide; heptad repeat region; transmembrane protein; gp41; HR1; HR2;  
XX infection.  
XX  
XX Human respiratory syncytial virus.  
XX  
XX  
XX Key Location/Qualifiers  
FT Modified-site 1  
FT /note= "N-terminal is substituted by Ac"  
FT Modified-site 34  
FT /note= "C-terminal amide"  
XX  
XX WO200164013-A2.  
XX  
XX 07-SEP-2001.  
XX  
XX 07-FEB-2001; 2001WO-US003988.  
XX  
XX 29-FEB-2000; 2000US-00515965.  
XX  
XX (TRIM-) TRIMERIS INC.  
XX  
XX Antczak JB, Delmedico MK, Erickson JB, Lambert DM, Sista P;  
XX WPI; 2001-514829/56.  
XX  
XX Heptad repeat region peptide analogs useful for inhibiting virus/cells  
PT fusion, useful for treating HIV and Respiratory Syncytial Virus  
PT infection.  
XX  
XX

XX Example; Page 40; 587pp; English.

XX The invention relates to isolated analogues of the heptad repeat region

XX peptides Dp178 and Dp107. Dp178 and Dp107 correspond to amino acids 638-

XX 673 (heptad repeat region HR2) and 558-595 (heptad repeat region HR1)

XX respectively, of HIV-1LAI transmembrane protein gp41. The HR1 and HR2

XX regions of proteins interact non-covalently with each other and/or with

XX peptides derived from them. This interaction is required for normal

XX infectivity of viruses such as RSV and HIV. The heptad repeat region

XX peptide analogues may be used to inhibit respiratory syncytial virus

XX (RSV) infection in a cell. They may also be used to inhibit HIV

XX infection. The present sequence is a peptide provided in the

XX specification

XX Sequence 34 AA;

Query Match 29.9%; Score 29; DB 4; Length 34;

Best Local Similarity 40.0%; Pred. No. 1.4e+03;

Matches 6; Conservative 2; Mismatches 7; Indels 0; Gaps 0;

QY 3 NMLNSKIAPKIVSQE 17

Db 13 NKXNGTDAVKLIKQE 27

RESULT 157

ABBO0366

ID ABB00366 standard; peptide; 34 AA.

AC ABB00366;

XX

DT 03-JAN-2002 (first entry)

XX

DE RSV F2 protein Dp178/107-like region peptide T371.

XX

KW Human immunodeficiency virus; HIV; respiratory syncytial virus; RSV;

KW virucide; heptad repeat region; transmembrane protein; gp41; HR1; HR2;

KW infection.

XX

OS Human respiratory syncytial virus.

XX

EH Key Location/Qualifiers

FT Modified-site 1 /note= "N-terminal is substituted by Ac"

FT Modified-site 34 /note= "C-terminal amide"

FT

XX WO200164013-A2.

PN

XX 07-SEP-2001.

PD

XX 07-FEB-2001; 2001WO-US003988.

PF

XX 29-FEB-2000; 2000US-00515965.

PR

XX (TRIM-) TRIMERIS INC.

PA

XX Antczak JB, Delmedico MK, Erickson JB, Lambert DM, Sista P;

PI WPI; 2001-514829/56.

DR

XX Heptad repeat region peptide analogs useful for inhibiting virus/cells

FT fusion, useful for treating HIV and Respiratory Syncytial Virus

PT infection.

PT

XX Example; Page 40; 587pp; English.

PS

XX The invention relates to isolated analogues of the heptad repeat region

XX peptides Dp178 and Dp107. Dp178 and Dp107 correspond to amino acids 638-

XX 673 (heptad repeat region HR2) and 558-595 (heptad repeat region HR1)

XX respectively, of HIV-1LAI transmembrane protein gp41. The HR1 and HR2

XX regions of proteins interact non-covalently with each other and/or with

CC peptides derived from them. This interaction is required for normal

CC infectivity of viruses such as RSV and HIV. The heptad repeat region

CC peptide analogues may be used to inhibit respiratory syncytial virus

CC (RSV) infection in a cell. They may also be used to inhibit HIV

CC infection. The present sequence is a peptide provided in the

CC specification

XX Sequence 34 AA;

Query Match 29.9%; Score 29; DB 4; Length 34;

Best Local Similarity 40.0%; Pred. No. 1.4e+03;

Matches 6; Conservative 2; Mismatches 7; Indels 0; Gaps 0;

QY 3 NMLNSKIAPKIVSQE 17

Db 15 NKXNGTDAVKLIKQE 29

RESULT 158

AAU12917

ID AAU12917 standard; peptide; 34 AA.

XX

AC AAU12917;

XX

DT 21-NOV-2001 (first entry)

XX

DE Dp178-like/Dp107-like peptide T-373.

XX

KW Anti-retroviral; Dp178-like; Dp107-like; transmembrane protein gp41;

KW antifusogenic; antiviral; HIV transmission; mutant; mutein.

KW

XX Human immunodeficiency virus 1; isolate LAI.

OS Synthetic.

XX

EH Key Location/Qualifiers

FT Modified-site 1 /note= "N-terminal is substituted by Ac"

FT Modified-site 34 /note= "C-terminal amide"

FT

XX WO200151673-A2.

PN

XX 19-JUL-2001.

PD

XX 05-JUL-2000; 2000WO-US035727.

PF

XX 09-JUL-1999; 99US-00350841.

PR

XX (TRIM-) TRIMERIS INC.

PA

XX Jeffs P, Lackey JW, Erickson JB, Lawless MK, Merucka G;

PI WPI; 2001-442157/47.

DR

XX Identifying a compound that inhibits the formation of or disrupts a

PT Dp107/Dp178 complex, especially compounds with antifusogenic, antiviral

PT or intracellular modulatory activity, by detecting the formation of a

PT Dp107/Dp178 complex.

PT

XX Disclosure; Page 60; 259pp; English.

PS

XX The present invention relates to peptides which exhibit anti-retroviral

CC activity. The peptides of the invention (AAU12559-AAU14009) comprise

CC Dp178-like and Dp107-like peptides. The Dp178 peptide corresponds to

CC amino acids 639-673 of the transmembrane protein gp41 from human

CC immunodeficiency virus 1 (HIV-1) isolate LAI. The Dp107 peptide

CC corresponds to amino acids 558-595 of gp41 from HIV-1LAI. The invention

CC also relates to a method of identifying compounds that inhibit the

CC formation of or disrupts a Dp107/Dp178 complex. The method comprises

CC detecting the formation of a Dp107/Dp178 complex, both in the presence or

CC absence of a test compound in a reaction mixture containing Dp107 and

CC Dp178 peptides. The method is useful for identifying compounds, including

CC small molecule compounds, which may themselves exhibit antifusogenic,



CC antiviral or intracellular modulatory activity. The DP178-like/DP107-like  
CC peptides are useful to inhibit human and non-human retroviral,  
CC particularly HIV, transmission to uninfected cells. The present sequence  
CC represents one of the DP178-like/DP107-like peptides of the invention

```

SQ      Sequence 34 AA;
      Query Match      29.9%;      Score 29;      DB 4;      Length 34;
      Best Local Similarity 40.0%;      Pred. No. 1.4e+03;
      Matches 6;      Conservative 2;      Mismatches 7;      Indels 0;      Gaps 0;

Qy      3      NHLNSKIAFKIVSQE 17
      |      |      |      |      |
Db      13      NKXNGTDAVKLIKOE 27

```

RESULT 159  
AAU12916  
ID AAU12916 standard; peptide; 34 AA.

AC	AAU12916;
XX	
XX	21-NOV-2001 (first entry)
XX	
XX	DP178-like/DP107-like peptide T-372.
XX	
XX	Anti-retroviral; DP178-like; transmembrane protein gp41;
KW	antifusogenic; antiviral; HIV transmission; mutant; mutesin.
XX	
XX	Human immunodeficiency virus 1; isolate LAI.
OS	Synthetic.
OS	

Key	Location/Qualifiers
PH	1
FT	/note= "N-terminal is substituted by Ac"
FT	34
FT	/note= "C-terminal amide"

PN	WO200151673-A2.
XX	
PD	19-JUL-2001.
XX	
XX	
PF	05-JUL-2000; 2000WO-US035727.
XX	
PR	09-JUL-1999; 98US-00350841.
XX	
XX	(TRIM-) TRIMERIS INC.
PA	
XX	
PI	Jeffs P, Lackey JW, Erickson JB, Lawless MK, Merutka G;
XX	
DR	WPI; 2001-442157/47.

Identifying a compound that inhibits the formation of or disrupts a DP107/DP178 complex, especially compounds with antifusogenic, antiviral or intracellular modulatory activity, by detecting the formation of a DP107/DP178 complex.

PS Disclosure; Page 60; 259pp; English.

XX

CC The present invention relates to peptides which exhibit anti-retroviral  
CC activity. The peptides of the invention (AAU12559-AAU14009) comprise  
CC DP178-like and DP107-like peptides. The DP178 peptide corresponds to  
CC amino acids 639-673 of the transmembrane protein gp41 from human  
CC immunodeficiency virus 1 (HIV-1) isolate LA1. The DP107 peptide  
CC corresponds to amino acids 558-595 of gp41 from HIV-1LA1. The invention  
CC also relates to a method of identifying compounds that inhibit the  
CC formation of or disrupts a DP107/DP178 complex. The method comprises  
CC detecting the formation of a DP107/DP178 complex, both in the presence or  
CC absence of a test compound, in a reaction mixture containing DP107 and  
CC DP178 peptides. The method is useful for identifying compounds, including  
CC small molecule compounds, which may themselves exhibit antiseogenic,  
CC antiviral or intracellular modulatory activity. The DP178-like/DP107-like  
CC peptides are useful to inhibit human and non-human retroviral.

CC particularly HIV, transmission to uninfected cells. The present sequence  
CC represents one of the DP178-like/DP107-like peptides of the invention  
XX  
SQ Sequence 34 AA;

```

Query Match      23.9%; Score 29; DB 4; Length 34;
Best Local Similarity 40.0%; Pred.No. 1.4e+03;
Matches 6; Conservative 2; Mismatches 7; Indels 0; Gaps 0
Qy      3  NHLNSKIAFIVSQF 17
Db      14  NKXNGTDAVKLIKOE 28

```

RESULT 160	
AAU12915	
ID	AAU12915 standard; peptide; 34 AA.
XX	
XX	
XX	AAU12915;
XX	
DT	21-NOV-2001 (first entry)
XX	
DE	DP178-like/DP107-like peptide T-371.
XX	
XX	
KW	Anti-retroviral; DP178-like; transmembrane protein gp41;
KW	antifusogenic; antiviral; HIV transmission; mutant; mutcin.
XX	
OS	Human immunodeficiency virus 1; isolate LAI.
OS	Synthetic.

Key	Location/Qualifiers
Modified-site	1
Modified-site	/note= "N-terminal is substituted by Ac"
Modified-site	34
Modified-site	/note= "C-terminal amide"

PN	WO200151673-A2.
XX	
XX	
PD	19-JUL-2001.
XX	
XX	
PF	05-JUL-2000; 2000WO-US035727.
XX	
PR	09-JUL-1999; 99US-00350841.
XX	
XX	(TRIM-) TRIMERIS INC.
PA	
XX	
PI	Jaffs P, Lackey JW, Erickson JB, Lawless MK, Merutka G;
XX	
DR	WPI; 2001-442157/47.

Identifying a compound that inhibits the formation of or disrupts a DP107/DP178 complex, especially compounds with antifusogenic, antiviral or intracellular modulatory activity, by detecting the formation of a DP107/DP178 complex.

PS Disclosure; Page 60; 259pp; English.

The present invention relates to peptides which exhibit anti-retroviral activity. The peptides of the invention (AAU12559-AAU14009) comprise DP178-like and DP107-like peptides. The DP178 peptide corresponds to amino acids 639-673 of the transmembrane protein gp41 from human immunodeficiency virus 1 (HIV-1) isolate LAI. The DP107 peptide corresponds to amino acids 558-595 of gp41 from HIV-1LAI. The invention also relates to a method of identifying compounds that inhibit the formation of or disrupts a DP107/DP178 complex. The method comprises detecting the formation of a DP107/DP178 complex, both in the presence or absence of a test compound, in a reaction mixture containing DP107 and DP178 peptides. The method is useful for identifying compounds, including small molecule compounds, which may themselves exhibit antifeedogenic, antiviral or intracellular modulatory activity. The DP178-like/DP107-like peptides are useful to inhibit human and non-human retroviral, particularly HIV, transmission to uninfected cells. The present sequence represents one of the DP178-like/DP107-like peptides of the invention

```
XX SQ Sequence 34 AA;
Query Match 29.9%; Score 29; DB 4; Length 34;
Best Local Similarity 40.0%; Pred. No. 1.4e+03;
Matches 6; Conservative 2; Mismatches 7; Indels 0; Gaps 0;

QY 3 NHLSKIAFKIVSQE 17
DB 15 NKXNGTDAVKLIKQE 29

RESULT 161
ADE01862
ID ADE01862 standard; peptide; 34 AA.
XX AC ADE01862;
XX DT 29-JAN-2004 (first entry)
XX DE Hybrid polypeptide pharmacokinetic enhancer peptide, SEQ ID No 369.
XX KW hybrid; enhancer; anti-fusogenic; antiviral; virucide; antidiabetic;
XX KW pharmacokinetic; fusogenic; insulin; diabetes.
XX OS Unidentified.
XX FH Key Location/Qualifiers
FT Modified-site 1 /note= "Residue is modified by acetyl group"
FT Modified-site 34 /note= "C-terminal amide"
XX PN US6348568-B1.
XX PD 19-FEB-2002.
XX PF 20-MAY-1999; 99US-00315304.
XX PR 20-MAY-1998; 98US-00082279.
XX PA (TRIM-) TRIMERIS INC.
XX PI Barney S, Guthrie KI, Merutka G, Anwer MK, Lambert DM;
XX WPI; 2002-424396/45.
XX New hybrid polypeptide for modulating fusogenic events for e.g. antiviral
PT activity, has enhancer peptide sequence derived from retroviral envelope
PT protein sequences linked to core polypeptide e.g. therapeutic protein.
XX PS Disclosure; SEQ ID NO 369; 70pp; English.
XX CC The invention relates to a novel hybrid polypeptide comprising an
CC enhancer peptide sequence linked to a core polypeptide. The enhancer
CC peptide sequence comprises WQWEQKI or WASLWEWF. The invention also
CC includes novel peptides that exhibit anti-fusogenic activity, antiviral
CC activity and/or ability to modulate intracellular processes. The novel
CC hybrid polypeptide has virucide and antidiabetic activity. The enhancer
CC peptide sequence enhances pharmacokinetic properties of any core
CC polypeptide, for example, a polypeptide useful for the treatment or
CC prevention of a disease, or an imaging agent useful for imaging
CC structures in vivo. The core polypeptides and hybrid polypeptides are
CC useful for modulating fusogenic events and exhibit antifusogenic or
CC antiviral activity. The novel hybrid polypeptide is useful for decreasing
CC viral infection and modulating intracellular processes involving coiled-
CC coil peptide interactions. The novel hybrid polypeptide comprises insulin
CC or its fragment, so the core polypeptide is useful for ameliorating the
CC symptoms of forms of diabetes. The novel hybrid polypeptide is also
CC useful as a part of prognosis for preventing disorders including fusion
CC events and viral infection that involves cell-cell and/or virus-cell
CC fusion, and for diagnosis and in vivo imaging methods. This sequence
CC represents an enhancer peptide of the invention.
```

```
XX SQ Sequence 34 AA;
Query Match 29.9%; Score 29; DB 5; Length 34;
Best Local Similarity 40.0%; Pred. No. 1.4e+03;
Matches 6; Conservative 2; Mismatches 7; Indels 0; Gaps 0;

QY 3 NHLSKIAFKIVSQE 17
DB 13 NKXNGTDAVKLIKQE 27

RESULT 162
ADE01860
ID ADE01860 standard; peptide; 34 AA.
XX AC ADE01860;
XX DT 29-JAN-2004 (first entry)
XX DE Hybrid polypeptide pharmacokinetic enhancer peptide, SEQ ID No 367.
XX KW hybrid; enhancer; anti-fusogenic; antiviral; virucide; antidiabetic;
XX KW pharmacokinetic; fusogenic; insulin; diabetes.
XX OS Unidentified.
XX FH Key Location/Qualifiers
FT Modified-site 1 /note= "Residue is modified by acetyl group"
FT Modified-site 34 /note= "C-terminal amide"
XX PN US6348568-B1.
XX PD 19-FEB-2002.
XX PF 20-MAY-1999; 99US-00315304.
XX PR 20-MAY-1998; 98US-00082279.
XX PA (TRIM-) TRIMERIS INC.
XX PI Barney S, Guthrie KI, Merutka G, Anwer MK, Lambert DM;
XX WPI; 2002-424396/45.
XX New hybrid polypeptide for modulating fusogenic events for e.g. antiviral
PT activity, has enhancer peptide sequence derived from retroviral envelope
PT protein sequences linked to core polypeptide e.g. therapeutic protein.
XX PS Disclosure; SEQ ID NO 367; 70pp; English.
XX CC The invention relates to a novel hybrid polypeptide comprising an
CC enhancer peptide sequence linked to a core polypeptide. The enhancer
CC peptide sequence comprises WQWEQKI or WASLWEWF. The invention also
CC includes novel peptides that exhibit anti-fusogenic activity, antiviral
CC activity and/or ability to modulate intracellular processes. The novel
CC hybrid polypeptide has virucide and antidiabetic activity. The enhancer
CC peptide sequence enhances pharmacokinetic properties of any core
CC polypeptide, for example, a polypeptide useful for the treatment or
CC prevention of a disease, or an imaging agent useful for imaging
CC structures in vivo. The core polypeptides and hybrid polypeptides are
CC useful for modulating fusogenic events and exhibit antifusogenic or
CC antiviral activity. The novel hybrid polypeptide is useful for decreasing
CC viral infection and modulating intracellular processes involving coiled-
CC coil peptide interactions. The novel hybrid polypeptide comprises insulin
CC or its fragment, so the core polypeptide is useful for ameliorating the
CC symptoms of forms of diabetes. The novel hybrid polypeptide is also
CC useful as a part of prognosis for preventing disorders including fusion
CC events and viral infection that involves cell-cell and/or virus-cell
CC fusion, and for diagnosis and in vivo imaging methods. This sequence
CC represents an enhancer peptide of the invention.
```

XX SQ Sequence 34 AA;  
 Query Match 29.9%; Score 29; DB 5; Length 34;  
 Best Local Similarity 40.0%; Pred. No. 1.4e+03;  
 Matches 6; Conservative 2; Mismatches 7; Indels 0; Gaps 0;  
 Qy 3 NMLNSKIAFKIVSQE 17  
 | | | | |  
 Db 15 NKXNGTDAVKLIKQE 29  
 | | | | |  
 RESULT 163  
 ABP29370  
 ID ABP29370 standard; protein; 37 AA.  
 XX AC  
 XX DT  
 XX DT 02-JUL-2002 (first entry)  
 XX DE Streptococcus polypeptide SEQ ID NO 7916.  
 XX KW Streptococcus; GAS; GBS; group B streptococcus; Streptococcus agalactiae;  
 XX KW group A streptococcus; Streptococcus pyogenes; antibacterial;  
 XX KW antiinflammatory; infection; vaccine; meningitis; gene therapy.  
 XX OS Streptococcus pyogenes.  
 XX PN WO200234771-A2.  
 XX PD 02-MAY-2002.  
 XX PF 29-OCT-2001; 2001WO-GB004789.  
 XX PR 27-OCT-2000; 2000GB-00026333.  
 XX PR 24-NOV-2000; 2000GB-00028727.  
 XX PR 07-MAR-2001; 2001GB-00005640.  
 XX PA (CHIR-) CHIRON SPA.  
 XX PA (GENO-) INST GENOMIC RES.  
 XX PI Telford J, Massignani V, Margarit Y Rosl, Grandi G, Fraser C;  
 XX PI Tettelin H;  
 XX PR WPI; 2002-352536/38.  
 XX DR N-PSDB; ABN70001.  
 XX PT New Streptococcus protein for the treatment or prevention of infection or  
 XX PT disease caused by Streptococcus bacteria, such as meningitis, and for  
 XX PT detecting a compound that binds to the protein.  
 XX PS Claim 1; Page 3921; 4525pp; English.  
 XX CC The invention relates to a protein (ABP25413-ABP30895) from group B  
 CC Streptococcus/GBS (Streptococcus agalactiae) or group A streptococcus/GAS  
 CC (Streptococcus pyogenes), comprising one of 5483 sequences (S1), given in  
 CC the specification. The proteins have antibacterial and antiinflammatory  
 CC activity. (I), nucleic acids encoding (I), ABN66044-ABN71526 and  
 CC antibodies that bind (I) are used in the manufacture of medicaments for  
 CC the treatment or prevention of infection or disease caused by  
 CC Streptococcus bacteria, particularly S. agalactiae and S. pyogenes.  
 CC Nucleic acids encoding (I) are used to detect Streptococcus in a  
 CC biological sample. (I) is used to determine whether a compound binds to  
 CC (I). A composition comprising (I) or a nucleic acid encoding (I), may be  
 CC used as a vaccine or diagnostic composition. The disease caused by  
 CC Streptococcus that is prevented or treated may be meningitis. Nucleic  
 CC acid encoding (I) may be used to recombinantly produce (I) and may be  
 CC used in gene therapy. Antibodies to (I) are used for affinity  
 CC chromatography, immunoassays, and distinguishing/identifying  
 XX Streptococcus proteins  
 XX SQ Sequence 37 AA;  
 Query Match 29.9%; Score 29; DB 5; Length 37;  
 Best Local Similarity 66.7%; Pred. No. 1.6e+03;  
 Matches 6; Conservative 2; Mismatches 1; Indels 0; Gaps 0;  
 Qy 8 KIAFKIVSQ 16  
 | | | | |  
 Db 12 KISLKIVAQ 20  
 | | | | |  
 RESULT 164  
 ADD90380  
 ID ADD90380 standard; protein; 37 AA.  
 XX AC  
 XX AC ADD90380;  
 XX DT  
 XX DT 29-JAN-2004 (first entry)  
 XX DE Novel human secreted protein seq id 11 protein feature seq id 195.  
 XX KW gene therapy; cytostatic; cancer; human; secreted protein.  
 XX OS Homo sapiens.  
 XX PN US2003199683-A1.  
 XX PD 23-OCT-2003.  
 XX PF 30-MAR-2001; 2001US-00820649.  
 XX PR 30-JUL-1997; 97US-0054209P.  
 XX PR 30-JUL-1997; 97US-0054211P.  
 XX PR 30-JUL-1997; 97US-0054212P.  
 XX PR 30-JUL-1997; 97US-0054213P.  
 XX PR 30-JUL-1997; 97US-0054214P.  
 XX PR 30-JUL-1997; 97US-0054215P.  
 XX PR 30-JUL-1997; 97US-0054217P.  
 XX PR 30-JUL-1997; 97US-0054218P.  
 XX PR 30-JUL-1997; 97US-0054234P.  
 XX PR 30-JUL-1997; 97US-0054236P.  
 XX PR 18-AUG-1997; 97US-0055968P.  
 XX PR 18-AUG-1997; 97US-0055969P.  
 XX PR 18-AUG-1997; 97US-0055972P.  
 XX PR 19-AUG-1997; 97US-0056534P.  
 XX PR 19-AUG-1997; 97US-0056543P.  
 XX PR 19-AUG-1997; 97US-0056554P.  
 XX PR 19-AUG-1997; 97US-0056561P.  
 XX PR 19-AUG-1997; 97US-0056727P.  
 XX PR 19-AUG-1997; 97US-0056729P.  
 XX PR 19-AUG-1997; 97US-0056730P.  
 XX PR 29-JUL-1998; 98WO-US015949.  
 XX PR 26-JAN-1999; 99US-00236557.  
 XX PR 21-SEP-2000; 2000US-00666987.  
 XX PA (RUBE/) RUBEN S M.  
 XX PA (FENG/) FENG P.  
 XX PA (LAFL/) LAFLEUR D W.  
 XX PA (MOOR/) MOORE P A.  
 XX PA (SHIY/) SHI Y.  
 XX PA (KYAW/) KYAW H.  
 XX PA (LIYV/) LI Y.  
 XX PA (ZENG/) ZENG Z.  
 XX PA (CART/) CARTER K C.  
 XX PA (ENDR/) ENDRESS G A.  
 XX PA (WEIY/) WEI Y.  
 XX PA (FANP/) FAN P.  
 XX PA (ROSE/) ROSEN C A.  
 XX PI Ruben SM, Feng P, Lafleur DW, Moore PA, Shi Y, Kyaw H, Li Y;  
 XX PI Zeng Z, Carter KP, Endress GA, Wei Y, Fan P, Rosen CA;  
 XX DR WPI; 2003-852813/79.  
 XX PT New nucleic acid molecule, useful for preparing a medicament for

PT preventing, treating or ameliorating a medical condition e.g., cancer.  
XX  
PS Disclosure; SEQ ID NO 195; 213pp; English.  
XX  
CC The invention describes novel isolated human nucleic acids. The nucleic acid is useful for preparing a medicament for preventing, treating or ameliorating a medical condition e.g., cancer, and in gene therapy. This is the amino acid sequence of polypeptide feature of a novel human secreted protein of the invention.  
XX  
SQ Sequence 37 AA;  
  
Query Match 29.9%; Score 29; DB 7; Length 37;  
Best Local Similarity 54.5%; Pred. No. 1.6e+03;  
Matches 6; Conservative 1; Mismatches 4; Indels 0; Gaps 0;  
  
QY 3 NHLNSKIAFKI 13  
DB 11 HHLKSKFHLKI 21  
  
RESULT 166  
ADG90199  
ID ADG90199 standard; peptide; 37 AA.  
AC ADG90199;  
XX  
DT 11-MAR-2004 (first entry)  
XX  
DE Human secreted protein gene 1 extra polypeptide #3.  
XX  
KW Secreted protein; gene therapy; neural disorder; immune system disorders; muscular disorder; reproductive disorder; gastrointestinal disorder;  
KW pulmonary disorder; cardiovascular disorder; renal disorder;  
KW proliferative disorder; cancer; systemic lupus erythematosus;  
KW rheumatoid arthritis; multiple sclerosis; thyroiditis; anaemia;  
KW Grave's disease; diabetes; hepatitis; asthma; allergy; nephritis;  
KW Parkinson's disease; Alzheimer's disease; atherosclerosis;  
KW myocardial infarction; AIDS; infection; human.  
XX  
OS Homo sapiens.  
XX  
PN US2003166541-A1.  
XX  
PD 04-SEP-2003.  
XX  
PF 04-JUN-2002; 2002US-00160162.  
XX  
PR 30-JUL-1997; 97US-0054203P.  
PR 30-JUL-1997; 97US-0054211P.  
PR 30-JUL-1997; 97US-0054212P.  
PR 30-JUL-1997; 97US-0054213P.  
PR 30-JUL-1997; 97US-0054214P.  
PR 30-JUL-1997; 97US-0054215P.  
PR 30-JUL-1997; 97US-0054217P.  
PR 30-JUL-1997; 97US-0054218P.  
PR 30-JUL-1997; 97US-0054234P.  
PR 30-JUL-1997; 97US-0054236P.  
PR 18-AUG-1997; 97US-0055968P.  
PR 18-AUG-1997; 97US-0055969P.  
PR 18-AUG-1997; 97US-0055972P.  
PR 19-AUG-1997; 97US-0056534P.  
PR 19-AUG-1997; 97US-0056543P.  
PR 19-AUG-1997; 97US-0056554P.  
PR 19-AUG-1997; 97US-0056561P.  
PR 19-AUG-1997; 97US-0056727P.  
PR 19-AUG-1997; 97US-0056729P.  
PR 19-AUG-1997; 97US-0056730P.  
PR 29-JUL-1998; 98WO-US015949.  
PR 26-JAN-1999; 99US-00236557.  
PR 05-JUN-2001; 2001US-0295558P.  
XX  
PA (HUMA-) HUMAN GENOME SCI INC.

XX Ruben SM, Feng P, Lafleur DW, Moore PA, Shi Y, Kyaw H, Li Y;  
PI Zeng Z, Carter KC, Endress GA, Wei Y, Fan P, Rosen CA;  
XX WPI; 2003-874923/81.  
DR  
XX Nucleic acids encoding 83 secreted polypeptides, useful for preventing, diagnosing and treating disorders related to their aberrant expression and activity.  
XX  
PS Disclosure; SEQ ID NO 195; 308pp; English.  
XX  
CC The invention relates to an isolated nucleic acid molecule encoding a secreted protein that is at least 95% identical to a polynucleotide fragment of any of the nucleotide sequences listed in table 1A of the specification, which is hybridisable to the nucleotide sequences, a polynucleotide encoding a polypeptide (or a polypeptide fragment, domain or epitope of any of the amino acid sequences) listed in table 1A of the specification, a polynucleotide which is an (allelic) variant of the nucleotide sequences listed in the specification, a polynucleotide which encodes a species homologue of the above amino acid sequences, a polynucleotide capable of hybridising under stringent conditions to any of the above polynucleotides, where the polynucleotide does not hybridise under stringent conditions to a nucleic acid molecule having a nucleotide sequence of only A or T residues. Also included are a recombinant vector comprising the above nucleic acid molecule, making a recombinant host cell comprising the above nucleic acid molecule, an isolated polypeptide comprising a sequence that is at least 95% identical to the polypeptide (or its fragment, domain, epitope, secreted form, (allelic) variant or homologue) encoded by the above nucleic acid molecule, an isolated antibody that binds specifically to the above polypeptide, a recombinant host cell produced by the above method and that expresses the above polypeptide, making an isolated polypeptide, preventing, treating or ameliorating a medical condition, diagnosing a pathological condition or a susceptibility to a pathological condition in a subject, identifying a binding partner to the above polypeptide, the gene corresponding to the cDNA sequence given in the specification, and identifying an activity in a biological assay. The nucleic acid molecule and polypeptide are useful in diagnosing, preventing, prognosing or treating diseases or disorders associated with aberrant expression and/or activity of the above polypeptide, such as neural disorders, immune system disorders, muscular disorders, reproductive disorders, gastrointestinal disorders, pulmonary disorders, cardiovascular disorders, renal disorders, proliferative disorders and/or cancers. In particular, these diseases are systemic lupus erythematosus, rheumatoid arthritis, multiple sclerosis, thyroiditis, anaemia, Grave's disease, diabetes, hepatitis, asthma, allergies, nephritis, Parkinson's disease, Alzheimer's disease, atherosclerosis, myocardial infarction, AIDS and infections. The methods may be used for identifying agonists and antagonists of the polynucleotide and polypeptide. The present sequence is a protein from one of the 83 disclosed secreted protein genes.  
XX  
SQ Sequence 37 AA;  
  
Query Match 29.9%; Score 29; DB 7; Length 37;  
Best Local Similarity 54.5%; Pred. No. 1.6e+03;  
Matches 6; Conservative 1; Mismatches 4; Indels 0; Gaps 0;  
  
QY 3 NHLNSKIAFKI 13  
DB 11 HHLKSKFHLKI 21  
  
RESULT 166  
ADL26415  
ID ADL26415 standard; peptide; 37 AA.  
XX  
AC ADL26415;  
XX  
DT 17-JUN-2004 (first entry)  
XX  
DE Synthetic peptide 1554 derived from a conserved region of HCV.  
XX

KW HCV; hepatitis C virus; virucide; vaccine; MHC; HLA;  
 KW major histocompatibility complex; human leukocyte antigen.  
 XX Synthetic.  
 XX WO2004024182-A2.  
 XX 25-MAR-2004.  
 XX 27-AUG-2003; 2003WO-EP009482.  
 XX 13-SEP-2002; 2002AT-00001376.  
 XX 27-FEB-2003; 2003WO-EP002005.  
 XX 11-JUL-2003; 2003EP-00450171.  
 XX (INTE-) INTERCELL AG.  
 XX Buschle M, Habel A, Klade C, Mattner F, Otava O, Vytvytska O;  
 XX Zauner W, Zinke S, Kirlappos H;  
 XX WPI; 2004-269899/25.  
 XX Isolating Hepatitis C Virus peptides (HVPs) which have a binding capacity  
 XX to a MHC/HLA molecule or a complex comprising the HCV-peptide and the  
 XX molecule by separating the complex from the HCV-peptides which do not  
 XX bind to the molecule.  
 XX Example 1; Page 32; 73pp; English.  
 XX The invention relates to a novel method for isolating Hepatitis C Virus  
 XX (HCV) peptides (HVPs). The method of the invention has virucide activity,  
 XX and may be useful in producing a vaccine. The method is useful for  
 XX isolating Hepatitis C Virus peptides (HVPs) which have a binding capacity  
 XX to a MHC/HLA molecule or a complex comprising the HCV-peptide and the  
 XX MHC/HLA molecule for preparing a vaccine against HCV infection. The  
 XX cells, a T cell clone or a T cell population or preparation is useful for  
 XX identifying heteroclitic epitopes or for preparing a composition for  
 XX treating HCV infection. The present sequence represents a synthetic  
 XX peptide derived from a conserved region of HCV.  
 XX Sequence 37 AA;

Query Match 29.9%; Score 29; DB 8; Length 37;  
 Best Local Similarity 55.6%; Pred. No. 1.6e+03;  
 Matches 5; Conservative 3; Mismatches 1; Indels 0; Gaps 0;  
 QY 9 IAFKIVSQE 17  
 Db :|||::|||  
 29 VAFKVMSE 37  
 RESULT 167  
 AAU87961  
 ID AAU87961 standard; peptide; 38 AA.  
 XX AAU87961;  
 AC  
 XX 05-JUN-2002 (first entry)  
 DT Human WW domain #14.  
 XX Human; PDZ domain; WW domain; rat; cow; mouse; fruitfly; protein therapy;  
 KW gene therapy; PDZ-mediated disease; inward potassium channel; WBP;  
 KW dimer inhibitor peptide; carboxylate binding loop.  
 XX Homo sapiens.  
 OS  
 XX WO200207751-A1.  
 XX 31-JAN-2002.  
 PD 24-JUL-2001; 2001WO-US023269.  
 XX

PR 25-JUL-2000; 2000US-0221215P.  
 PR 28-NOV-2000; 2000US-00723810.  
 XX (AXCE-) AXCELL BIOSCIENCES CORP.  
 XX Herrero J, Pirozzi G, Uveges A;  
 PI WPI; 2002-195842/25.  
 DR  
 XX Methods for identifying polypeptides comprising PDZ domains, the  
 PT polypeptides and their encoding nucleic acids, useful for the diagnosis  
 PT and treatment of PDZ related disorders.  
 XX  
 XX Disclosure; Fig 20; 225pp; English.  
 PS  
 CC The invention relates to methods for identifying polypeptides comprising  
 CC PDZ domains, and their encoding nucleic acids. The sequences are used to  
 CC identify modulators of their expression, function and activity, for use  
 CC in the diagnosis and treatment of PDZ related disorders. Antibodies  
 CC against the proteins and cells that produce them may be used for the  
 CC treatment of PDZ-mediated disease states. Sequences AAU87843-AAU87974  
 CC represent proteins containing PDZ domains, fragments of these proteins  
 CC and other related peptides used in the methods of the invention  
 XX Sequence 38 AA;  
 SQ  
 Query Match 29.9%; Score 29; DB 5; Length 38;  
 Best Local Similarity 50.0%; Pred. No. 1.6e+03;  
 Matches 5; Conservative 2; Mismatches 3; Indels 0; Gaps 0;  
 QY 3 NHLNKKIAFK 12  
 Db :|||::|||  
 21 DHLNKKIQFE 30  
 RESULT 168  
 AAU87961  
 ID AAU87961 standard; peptide; 38 AA.  
 XX AAU87961;  
 AC  
 XX 26-JUL-2002 (first entry)  
 DT  
 XX BS203 consensus sequence antigenic peptide #4.  
 DE  
 XX BS203 protein; therapy; breast disease; tumour; metastasis; cancer.  
 KW  
 XX Unidentified.  
 OS  
 XX US2002042049-A1.  
 PN  
 XX 11-APR-2002.  
 PD  
 XX 16-FEB-1999; 99US-00250883.  
 PF  
 XX 08-JUL-1997; 97US-00889316.  
 PR  
 XX (RUSS/) RUSSELL J C.  
 PA (COLP/) COLPITTS T L.  
 PA  
 XX Russell JC, Colpitts TL;  
 PI WPI; 2002-315123/35.  
 DR  
 XX Detecting a target BS203 polynucleotide in a test sample, is useful for  
 PT diagnosing diseases of the breast, specifically breast cancer.  
 PT  
 XX Claim 7; Page 39; 45pp; English.  
 PS  
 XX The invention relates to a set of contiguous and partially overlapping  
 CC cDNA sequences and polypeptides encoded thereby, designated as BS203. The  
 CC invention also provides antibodies which specifically bind to BS203-  
 CC encoded polypeptide or protein, and agonists or inhibitors which prevent

CC action of the tissue-specific BS203 polypeptide, are useful for the  
 CC therapeutic treatment of breast disease, tumours or metastases. The  
 CC sequences of the invention are useful for detecting, diagnosing, staging,  
 CC monitoring, prognosticating, in vivo imaging, preventing or treating, or  
 CC determining the predisposition of an individual to diseases and  
 CC conditions of the breast, such as breast cancer. The present sequence is  
 CC BS203 consensus sequence antigenic peptide

XX Sequence 38 AA;

Query Match 29.9%; Score 29; DB 5; Length 38;  
 Best Local Similarity 83.3%; Pred. No. 1.6e+03;  
 Matches 5; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 4 HNSKI 9  
 DB 18 HNSKL 23

RESULT 169  
 AAW20028  
 ID AAW20028 standard; protein; 39 AA.

AC AAW20028;  
 DT 25-MAR-2003 (revised)  
 DT 18-SEP-1997 (first entry)  
 XX Human acidic fibroblast growth factor exon 3 product.  
 DE FGF; fibroblast growth factor; basic; acidic; wound healing;  
 XX neurodegenerative disease; Parkinson's; Alzheimer's disease;  
 KW bone fracture; biologically active; embolism; bacteriophage.  
 XX

OS Homo sapiens.

XX US5604293-A.  
 PN 18-FEB-1997.  
 PD 01-APR-1994; 94US-00221462.  
 XX 12-SEP-1985; 85US-00775521.  
 PR 16-DEC-1985; 85US-00809163.  
 PR 30-MAY-1986; 86US-00869382.  
 PR 15-MAY-1987; 87US-00050786.  
 PR 30-MAR-1992; 92US-00860688.

PA (SCIO-) SCIOS INC.

XX Fiddes JC, Abraham JA;

PI WPI; 1997-234676/21.  
 DR N-PSDB; AAT71234.

XX New high purity, recombinant human basic fibroblast growth factor - for  
 PT promoting wound healing and treating neurodegenerative diseases,  
 PT suitable for production on large scale.

XX Example 3; Fig 2c; 34pp; English.

XX AAW20028 is the exon 3 product of human acidic fibroblast growth factor  
 CC (aFGF) derived from bacteriophage lambda-HAG-3. DNA encoding this product  
 CC was used to produce a recombinant aFGF protein. FGF is used to promote  
 CC healing of wounds, bone fractures, damaged myocardial tissue etc. and,  
 CC since it increases neuronal survival and promotes neurite outgrowth, may  
 CC also be used in treatment of neurological disorders such as Alzheimer's  
 CC and Parkinson's diseases. bFGF (basic FGF) may also be used for detection  
 CC of specific inhibitors; for treatment of cell cultures in vitro before  
 CC transplant and for inducing release of tissue plasminogen activator or  
 CC collagenase, e.g. for treatment of a chronic tendency to form embolism.  
 CC Recombinant FGFs can be produced on a large scale. (Updated on 25-MAR-  
 CC 2003 to correct PF field.)

XX SQ Sequence 39 AA;

Query Match 29.9%; Score 29; DB 2; Length 39;  
 Best Local Similarity 50.0%; Pred. No. 1.7e+03;  
 Matches 6; Conservative 2; Mismatches 4; Indels 0; Gaps 0;

QY 1 EPNHLSKIAPK 12  
 DB 14 EENHYNTYISKK 25

RESULT 170  
 AAB53950  
 ID AAB53950 standard; protein; 39 AA.

XX AC AAB53950;

DT 09-MAR-2001 (first entry)

XX Human colon cancer antigen protein sequence SEQ ID NO:1490.

XX Human; colon cancer; colon cancer antigen; diagnosis; detection;  
 KW identification; cytostatic; cardioactive; neuroprotective; vulnery;  
 KW immunomodulatory; muscular; gynaecological; gastrointestinal;  
 KW nephrotropic; antiinfective; antibacterial; gene therapy; wound;  
 KW neural disorder; immune system disorder; muscular disorder;  
 KW reproductive disorder; gastrointestinal disorder; renal disorder;  
 KW infectious disease; cardiovascular disorder.

XX Homo sapiens.

XX WO200055351-A1.

XX PD 21-SEP-2000.

XX PF 08-MAR-2000; 2000WO-US005883.

XX PR 12-MAR-1999; 99US-0124270P.

XX PA (HUMA-) HUMAN GENOME SCI INC.

XX Rosen CA, Ruben SM;

DR WPI; 2000-587534/55.  
 DR N-PSDB; AAC98707.

XX Colon cancer associated gene sequences, referred to as colon cancer  
 PT antigens, useful for the treatment, prevention, and diagnosis of colon  
 PT disorders such as colon cancer.

XX Claim 11; Page 2050; 2104pp; English.

XX AAC97991 to AAC98763 encode the human colon cancer associated proteins,  
 CC called human colon cancer antigens, given in AAB53234 to AAB54006. The  
 CC human colon cancer antigens can have cytostatic, cardioactive, muscular;  
 CC neuroprotective, immunomodulatory, gynaecological, gastrointestinal,  
 CC vulnery, nephrotropic, antiinfective and antibacterial activities, and  
 CC can be used in gene therapy. The colon cancer antigen polynucleotides,  
 CC proteins and antibodies to the proteins are useful for the prevention,  
 CC treatment and diagnosis of colon disorders, such as colon cancer. The  
 CC polynucleotides may be used in diagnostics and research, such as for  
 CC chromosome identification, and as hybridisation probes. The proteins may  
 CC also be used to prevent diseases such as neural disorders, immune system  
 CC disorders, muscular disorders, reproductive disorders, gastrointestinal  
 CC disorders, wounds, renal disorders, infectious diseases, and  
 CC cardiovascular disorders. AAC98764 to AAC98772 and AAB54007 represent  
 CC sequences used in the exemplification of the present invention

XX SQ Sequence 39 AA;

Query Match 29.9%; Score 29; DB 3; Length 39;  
 Best Local Similarity 37.5%; Pred. No. 1.7e+03;

Matches 6; Conservative 3; Mismatches 7; Indels 0; Gaps 0;

Qy 3 NHLNSKIAFKIVSQEP 18  
 Db 14 NLMTGRHSFKTYSQXP 29

RESULT 171  
 ABO55502  
 ID ABO55502 standard; protein; 40 AA.  
 XX  
 AC ABO55502;  
 XX  
 DT 29-JUL-2004 (first entry)  
 XX  
 DE Human genome derived single exon protein #1736.  
 XX  
 KW Human; gene expression; single exon probe; microarray;  
 KW alternative splicing event; genomic alteration.  
 XX  
 OS Homo sapiens.  
 XX  
 PN US2003194704-A1.  
 XX  
 PD 16-OCT-2003.  
 XX  
 XX 03-APR-2002; 2002US-00029386.  
 XX  
 PR 03-APR-2002; 2002US-00029386.  
 XX  
 PA (PENN/) PENN S G.  
 PA (RANK/) RANK D R.  
 PA (HANZ/) HANZEL D K.  
 XX  
 PI Penn SG, Rank DR, Hanzel DK;  
 XX  
 XX WPI; 2004-119264/12.  
 DR  
 XX  
 XX New human genome-derived single exon nucleic acid probes useful for human  
 PT gene expression analysis, for identifying or characterizing alternative  
 PT splicing events, for assessing genomic alterations or as tools for  
 PT surveying tissues.  
 XX  
 PS Claim 45; SEQ ID NO 29136; 80pp; English.  
 XX  
 CC The invention relates to a nucleic acid probe for measuring human gene  
 CC expression, comprising any of the 27,400 fully defined nucleotide  
 CC sequences in the specification, or their complements or fragments, and  
 CC encoding at least 8 amino acids of any of the 688 amino acid sequences  
 CC fully defined in the specification. The probe is a single exon probe that  
 CC hybridises under high stringency conditions to a nucleic acid molecule  
 CC expressed in human cells or tissues. Also included are a spatially-  
 CC addressable set of single exon nucleic acid probes for measuring human  
 CC gene expression (comprising a plurality of single exon nucleic acid  
 CC probes cited above, where each of the plurality of probes is separately  
 CC and addressably isolatable or amplifiable from the plurality), a single  
 CC exon microarray for measuring human gene expression, a method of  
 CC measuring human gene expression, a vector comprising the single exon  
 CC probe cited above, an ORF-encoded peptide comprising at least 8  
 CC contiguous amino acids of any of the above-mentioned amino acid  
 CC sequences (optionally with conservative amino acid substitutions), an  
 CC isolated antibody that binds specifically to a peptide cited above,  
 CC methods of selling and/or licensing single exon probes or microarrays to  
 CC a customer desiring to measure gene expression, a method of providing  
 CC human gene expression data by subscription, and a computer-readable  
 CC storage medium which contains a database having a plurality of records  
 CC (each record including data on the expression of a single exon probe  
 CC cited above). The probe, methods and apparatus are useful in gene  
 CC expression analysis. The probes may be used as tools for surveying  
 CC tissues to detect the presence of expressed messages that contain their  
 CC specific exon, or in constructing genome-derived single exon microarrays.  
 CC In addition, the probes are used in identifying and characterising  
 CC alternative splicing events, in detecting and characterising gross

CC alterations in the genomic locus that includes their exon, in assessing  
 CC smaller genomic alterations, in priming the synthesis of nucleic acids,  
 CC or in expressing the ORF-encoded peptide. The present sequence is a human  
 CC single exon probe protein of the invention. Note: The sequence data for  
 CC this patent did not form part of the printed specification, but was  
 CC obtained in electronic format directly from USPTO at  
 CC seqdata.uspto.gov/sequence.html?DocID=20030194704  
 XX  
 SQ Sequence 40 AA;

Query Match 29.9%; Score 29; DB 8; Length 40;  
 Best Local Similarity 54.5%; Pred No. 1.7e+03;  
 Matches 6; Conservative 1; Mismatches 4; Indels 0; Gaps 0;

Qy 3 NHLNSKIAFKI 13  
 Db 27 NHLNNTIVSHI 37

RESULT 172  
 ABO59597  
 ID ABO59597 standard; protein; 41 AA.  
 XX  
 AC ABO59597;  
 XX  
 DT 29-JUL-2004 (first entry)  
 XX  
 DE Human genome derived single exon protein #5831.  
 XX  
 KW Human; gene expression; single exon probe; microarray;  
 KW alternative splicing event; genomic alteration.  
 XX  
 OS Homo sapiens.  
 XX  
 PN US2003194704-A1.  
 XX  
 PD 16-OCT-2003.  
 XX  
 XX 03-APR-2002; 2002US-00029386.  
 XX  
 PR 03-APR-2002; 2002US-00029386.  
 XX  
 PA (PENN/) PENN S G.  
 PA (RANK/) RANK D R.  
 PA (HANZ/) HANZEL D K.  
 XX  
 PI Penn SG, Rank DR, Hanzel DK;  
 XX  
 XX WPI; 2004-119264/12.  
 DR  
 XX  
 XX New human genome-derived single exon nucleic acid probes useful for human  
 PT gene expression analysis, for identifying or characterizing alternative  
 PT splicing events, for assessing genomic alterations or as tools for  
 PT surveying tissues.  
 XX  
 PS Claim 45; SEQ ID NO 32231; 80pp; English.  
 XX  
 CC The invention relates to a nucleic acid probe for measuring human gene  
 CC expression, comprising any of the 27,400 fully defined nucleotide  
 CC sequences in the specification, or their complements or fragments, and  
 CC encoding at least 8 amino acids of any of the 688 amino acid sequences  
 CC fully defined in the specification. The probe is a single exon probe that  
 CC hybridises under high stringency conditions to a nucleic acid molecule  
 CC expressed in human cells or tissues. Also included are a spatially-  
 CC addressable set of single exon nucleic acid probes for measuring human  
 CC gene expression (comprising a plurality of single exon nucleic acid  
 CC probes cited above, where each of the plurality of probes is separately  
 CC and addressably isolatable or amplifiable from the plurality), a single  
 CC exon microarray for measuring human gene expression, a method of  
 CC measuring human gene expression, a vector comprising the single exon  
 CC probe cited above, an ORF-encoded peptide comprising at least 8  
 CC contiguous amino acids of any of the above-mentioned amino acid  
 CC sequences (optionally with conservative amino acid substitutions), an  
 CC isolated antibody that binds specifically to a peptide cited above,  
 CC methods of selling and/or licensing single exon probes or microarrays to  
 CC a customer desiring to measure gene expression, a method of providing  
 CC human gene expression data by subscription, and a computer-readable  
 CC storage medium which contains a database having a plurality of records  
 CC (each record including data on the expression of a single exon probe  
 CC cited above). The probe, methods and apparatus are useful in gene  
 CC expression analysis. The probes may be used as tools for surveying  
 CC tissues to detect the presence of expressed messages that contain their  
 CC specific exon, or in constructing genome-derived single exon microarrays.  
 CC In addition, the probes are used in identifying and characterising  
 CC alternative splicing events, in detecting and characterising gross

CC isolated antibody that binds specifically to a peptide cited above.  
CC methods of selling and/or licensing single exon probes or microarrays to  
CC a customer desiring to measure gene expression, a method of providing  
CC human gene expression data by subscription, and a computer-readable  
CC storage medium which contains a database having a plurality of records  
CC (each record including data on the expression of a single exon probe  
CC cited above. The probe, methods and apparatus are useful in gene  
CC expression analysis. The probes may be used as tools for surveying  
CC tissues to detect the presence of expressed messages that contain their  
CC specific exon, or in constructing genome-derived single exon microarrays.  
CC In addition, the probes are used in identifying and characterizing  
CC alternative splicing events, in detecting and characterising gross  
CC alterations in the genomic locus that includes their exon, in assessing  
CC smaller genomic alterations, in priming the synthesis of nucleic acids,  
CC or in expressing the ORF-encoded peptide. The present sequence is a human  
CC single exon probe protein of the invention. Note: The sequence data for  
CC this patent did not form part of the printed specification, but was  
CC obtained in electronic format directly from USPTO at  
CC seqdata.uspto.gov/sequence.html?DocID=20030194704  
XX  
XX  
SQ Sequence 41 AA;

Query Match 29.9%; Score 29; DB 8; Length 41;  
Best Local Similarity 83.3%; Pred. No. 1.8e+03;  
Matches 5; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 4 HNSKI 9  
Db 35 HNSK 40  
|||||

RESULT 173  
AAB64605  
ID AAB64605 standard; protein; 42 AA.  
XX  
AC AAB64605;  
XX  
DT 22-MAR-2001 (first entry)  
XX  
DE Human secreted protein BLAST search protein SEQ ID NO: 115.  
XX  
KW Cytostatic; immunosuppressive; nootropic; neuroprotective; antiviral;  
KW anti-allergic; hepatotropic; antidiabetic; anti-inflammatory; anti-ulcer;  
KW vulnerary; anticonvulsant; antibacterial; antifungal; antiparasitic;  
KW cardiant; gene therapy; cancer; immune disorder; cardiovascular disorder;  
KW neurological disease; infection; human; secreted protein.  
XX  
OS Homo sapiens.  
XX  
PN WO200077197-A1.  
XX  
PD 21-DEC-2000.  
XX  
PF 01-JUN-2000; 2000WO-US014934.  
XX  
PR 11-JUN-1999; 99US-0138599P.  
XX  
XX (HUMA-) HUMAN GENOME SCI INC.  
PA (ROSE/) ROSEN C A.  
XX  
XX Rosen CA, Ruben SM, Komatsoulis GA;  
XX WPI; 2001-032312/04.  
XX  
XX Isolated nucleic acid molecule encoding a human secreted protein is used  
PT in preventing, treating or ameliorating a medical condition.  
XX  
XX Disclosure; Page 512; 558pp; English.  
XX  
XX The invention relates to the isolation of genes AAB32757-F32803 encoding  
CC the human secreted proteins AAB64549-B64594. The sequence is a search  
CC result from a BLASTX homology search. The genes and proteins are useful  
CC for preventing, ameliorating or treating medical conditions, e.g. by

CC protein or gene therapy. The genes are isolated from a range of human  
CC tissues disclosed in the specification. The nucleic acids, proteins,  
CC antibodies and (ant)agonists are useful in the diagnosis, treatment and  
CC prevention of: (a) cancer, e.g. breast and ovarian cancer, and other  
CC cancers of the adrenal gland, bone, bone marrow, breast, gastrointestinal  
CC tract, liver, lung, or urogenital; (b) immune disorders e.g. Addison's  
CC disease, allergies, autoimmune haemolytic anaemia, autoimmune  
CC thyroiditis, diabetes mellitus, Crohn's disease, multiple sclerosis,  
CC rheumatoid arthritis and ulcerative colitis; (c) cardiovascular disorders  
CC such as myocardial ischaemias; (d) wound healing; (e) neurological  
CC diseases e.g. cerebral anoxia and epilepsy; and (f) infectious diseases  
CC such as viral, bacterial, fungal and parasitic infections  
XX  
SQ Sequence 42 AA;

Query Match 29.9%; Score 29; DB 4; Length 42;  
Best Local Similarity 71.4%; Pred. No. 1.8e+03;  
Matches 5; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

QY 1 EPNHLS 7  
Db 19 QENHLS 25  
:|||||

RESULT 174  
AAG09091  
ID AAG09091 standard; protein; 43 AA.  
XX  
AC AAG09091;  
XX  
DT 17-OCT-2000 (first entry)  
XX  
DE Arabidopsis thaliana protein fragment SEQ ID NO: 6886.  
XX  
KW Protein identification; signal transduction pathway; metabolic pathway;  
KW hybridisation assay; genetic mapping; gene expression control; promoter;  
KW termination sequence.  
XX  
OS Arabidopsis thaliana.  
XX  
PN EP1033405-A2.  
XX  
XX 06-SEP-2000.  
XX  
PF 25-FEB-2000; 2000EP-00301439.  
XX  
PR 25-FEB-1999; 99US-0121825P.  
PR 05-MAR-1999; 99US-0123180P.  
PR 09-MAR-1999; 99US-0123548P.  
PR 23-MAR-1999; 99US-0125788P.  
PR 25-MAR-1999; 99US-0126264P.  
PR 29-MAR-1999; 99US-0126785P.  
PR 01-APR-1999; 99US-0127462P.  
PR 06-APR-1999; 99US-0128234P.  
PR 08-APR-1999; 99US-0128714P.  
PR 16-APR-1999; 99US-0129845P.  
PR 19-APR-1999; 99US-0130077P.  
PR 21-APR-1999; 99US-0130449P.  
PR 23-APR-1999; 99US-0130510P.  
PR 23-APR-1999; 99US-0130891P.  
PR 28-APR-1999; 99US-0131449P.  
PR 30-APR-1999; 99US-0132048P.  
PR 30-APR-1999; 99US-0132407P.  
PR 04-MAY-1999; 99US-0132484P.  
PR 05-MAY-1999; 99US-0132485P.  
PR 06-MAY-1999; 99US-0132486P.  
PR 06-MAY-1999; 99US-0132487P.  
PR 07-MAY-1999; 99US-0132863P.  
PR 11-MAY-1999; 99US-0134256P.  
PR 14-MAY-1999; 99US-0134218P.  
PR 14-MAY-1999; 99US-0134219P.  
PR 14-MAY-1999; 99US-0134221P.  
PR 14-MAY-1999; 99US-0134370P.



PR 18-MAY-1999; 99US-0134768P.  
PR 19-MAY-1999; 99US-0134941P.  
PR 20-MAY-1999; 99US-0135124P.  
PR 21-MAY-1999; 99US-0135353P.  
PR 24-MAY-1999; 99US-0135622P.  
PR 25-MAY-1999; 99US-0136021P.  
PR 27-MAY-1999; 99US-0136392P.  
PR 28-MAY-1999; 99US-0136782P.  
PR 01-JUN-1999; 99US-0137222P.  
PR 03-JUN-1999; 99US-0137528P.  
PR 04-JUN-1999; 99US-0137502P.  
PR 07-JUN-1999; 99US-0137724P.  
PR 08-JUN-1999; 99US-0138094P.  
PR 10-JUN-1999; 99US-0138540P.  
PR 10-JUN-1999; 99US-0138847P.  
PR 14-JUN-1999; 99US-0139119P.  
PR 16-JUN-1999; 99US-0139452P.  
PR 16-JUN-1999; 99US-0139453P.  
PR 17-JUN-1999; 99US-0139492P.  
PR 18-JUN-1999; 99US-0139454P.  
PR 18-JUN-1999; 99US-0139455P.  
PR 18-JUN-1999; 99US-0139456P.  
PR 18-JUN-1999; 99US-0139457P.  
PR 18-JUN-1999; 99US-0139458P.  
PR 18-JUN-1999; 99US-0139459P.  
PR 18-JUN-1999; 99US-0139460P.  
PR 18-JUN-1999; 99US-0139461P.  
PR 18-JUN-1999; 99US-0139462P.  
PR 18-JUN-1999; 99US-0139463P.  
PR 18-JUN-1999; 99US-0139750P.  
PR 21-JUN-1999; 99US-0139763P.  
PR 22-JUN-1999; 99US-0139899P.  
PR 23-JUN-1999; 99US-0140333P.  
PR 23-JUN-1999; 99US-0140354P.  
PR 24-JUN-1999; 99US-0140695P.  
PR 28-JUN-1999; 99US-0140823P.  
PR 29-JUN-1999; 99US-0140991P.  
PR 30-JUN-1999; 99US-0141287P.  
PR 01-JUL-1999; 99US-0141842P.  
PR 01-JUL-1999; 99US-0142154P.  
PR 02-JUL-1999; 99US-0142055P.  
PR 06-JUL-1999; 99US-0142330P.  
PR 08-JUL-1999; 99US-0142803P.  
PR 09-JUL-1999; 99US-0142920P.  
PR 12-JUL-1999; 99US-0142977P.  
PR 13-JUL-1999; 99US-0143342P.  
PR 14-JUL-1999; 99US-0143624P.  
PR 15-JUL-1999; 99US-0144005P.  
PR 16-JUL-1999; 99US-0144085P.  
PR 16-JUL-1999; 99US-0144086P.  
PR 19-JUL-1999; 99US-0144325P.  
PR 19-JUL-1999; 99US-0144331P.  
PR 19-JUL-1999; 99US-0144332P.  
PR 19-JUL-1999; 99US-0144333P.  
PR 19-JUL-1999; 99US-0144334P.  
PR 19-JUL-1999; 99US-0144335P.  
PR 20-JUL-1999; 99US-0144352P.  
PR 20-JUL-1999; 99US-0144632P.  
PR 20-JUL-1999; 99US-0144884P.  
PR 21-JUL-1999; 99US-0144814P.  
PR 21-JUL-1999; 99US-0145086P.  
PR 21-JUL-1999; 99US-0145088P.  
PR 22-JUL-1999; 99US-0145085P.  
PR 22-JUL-1999; 99US-0145087P.  
PR 22-JUL-1999; 99US-0145089P.  
PR 23-JUL-1999; 99US-0145192P.  
PR 23-JUL-1999; 99US-0145145P.  
PR 23-JUL-1999; 99US-0145218P.  
PR 23-JUL-1999; 99US-0145224P.  
PR 26-JUL-1999; 99US-0145276P.  
PR 27-JUL-1999; 99US-0145913P.  
PR 27-JUL-1999; 99US-0145918P.  
PR 27-JUL-1999; 99US-0145919P.  
PR 28-JUL-1999; 99US-0145951P.  
PR 02-AUG-1999; 99US-0146386P.  
PR 02-AUG-1999; 99US-0146387P.  
PR 02-AUG-1999; 99US-0146388P.  
PR 03-AUG-1999; 99US-0146389P.  
PR 04-AUG-1999; 99US-0147038P.  
PR 04-AUG-1999; 99US-0147284P.  
PR 05-AUG-1999; 99US-0147302P.  
PR 05-AUG-1999; 99US-0147192P.  
PR 06-AUG-1999; 99US-0147260P.  
PR 06-AUG-1999; 99US-0147303P.  
PR 06-AUG-1999; 99US-0147416P.  
PR 09-AUG-1999; 99US-0147433P.  
PR 09-AUG-1999; 99US-0147935P.  
PR 10-AUG-1999; 99US-0148171P.  
PR 11-AUG-1999; 99US-0148319P.  
PR 12-AUG-1999; 99US-0148341P.  
PR 13-AUG-1999; 99US-0148565P.  
PR 13-AUG-1999; 99US-0148684P.  
PR 16-AUG-1999; 99US-0149368P.  
PR 17-AUG-1999; 99US-0149175P.  
PR 18-AUG-1999; 99US-0149426P.  
PR 20-AUG-1999; 99US-0149722P.  
PR 20-AUG-1999; 99US-0149723P.  
PR 23-AUG-1999; 99US-0149929P.  
PR 23-AUG-1999; 99US-0149902P.  
PR 23-AUG-1999; 99US-0150566P.  
PR 25-AUG-1999; 99US-0149930P.  
PR 26-AUG-1999; 99US-0150884P.  
PR 27-AUG-1999; 99US-0151065P.  
PR 27-AUG-1999; 99US-0151066P.  
PR 27-AUG-1999; 99US-0151080P.  
PR 30-AUG-1999; 99US-0151303P.  
PR 31-AUG-1999; 99US-0151438P.  
PR 01-SEP-1999; 99US-0151930P.  
PR 07-SEP-1999; 99US-0152363P.  
PR 10-SEP-1999; 99US-0153070P.  
PR 13-SEP-1999; 99US-0153758P.  
PR 15-SEP-1999; 99US-0154018P.  
PR 16-SEP-1999; 99US-0154039P.  
PR 20-SEP-1999; 99US-0154779P.  
PR 22-SEP-1999; 99US-0155139P.  
PR 23-SEP-1999; 99US-0155486P.  
PR 28-SEP-1999; 99US-0155659P.  
PR 29-SEP-1999; 99US-0156458P.  
PR 04-OCT-1999; 99US-0157117P.  
PR 05-OCT-1999; 99US-0157753P.  
PR 06-OCT-1999; 99US-0157865P.  
PR 07-OCT-1999; 99US-0158029P.  
PR 08-OCT-1999; 99US-0158232P.  
PR 12-OCT-1999; 99US-0158369P.  
PR 13-OCT-1999; 99US-0159293P.  
PR 13-OCT-1999; 99US-0159294P.  
PR 14-OCT-1999; 99US-0159295P.  
PR 14-OCT-1999; 99US-0159329P.  
PR 14-OCT-1999; 99US-0159330P.  
PR 14-OCT-1999; 99US-0159331P.  
PR 14-OCT-1999; 99US-0159637P.  
PR 18-OCT-1999; 99US-0159638P.  
PR 21-OCT-1999; 99US-0159584P.  
PR 21-OCT-1999; 99US-0160741P.  
PR 21-OCT-1999; 99US-0160767P.  
PR 21-OCT-1999; 99US-0160768P.  
PR 21-OCT-1999; 99US-0160770P.  
PR 21-OCT-1999; 99US-0160814P.  
PR 21-OCT-1999; 99US-0160815P.  
PR 22-OCT-1999; 99US-0160980P.  
PR 22-OCT-1999; 99US-0160981P.  
PR 22-OCT-1999; 99US-0160989P.  
PR 25-OCT-1999; 99US-0161404P.  
PR 25-OCT-1999; 99US-0161405P.  
PR 25-OCT-1999; 99US-0161406P.

XX The invention discloses isolated colon specific nucleic acids (CSNAs) and  
CC the polypeptides (CSPs) that they encode. The colon specific nucleic  
CC acids and polypeptides are useful as vaccines. The colon specific nucleic  
CC acid and polypeptides are also useful for diagnosing and monitoring the  
CC presence and metastases of colon cancer in a patient. The antibody that  
CC specifically binds to the colon specific polypeptide is useful for  
CC determining the presence of a colon specific polypeptide in a sample, as well  
CC as for treating a patient with cancer, particularly by inducing an immune  
CC response against the colon cancer cell expressing the colon specific  
CC nucleic acid molecule or polypeptide. In particular, these colon specific  
CC genes and proteins are useful for identifying, diagnosing, monitoring,  
CC staging, imaging and treating colon cancer (e.g. colorectal cancer) and  
CC non-cancerous disease states in the colon. These are also useful in gene  
CC therapy, production of transgenic animals and cells and in the production  
CC of engineered colon tissue for treatment and research. The sequences  
CC presented in ABG90928-ABG90963 are the human CSPs encoded by the CSNAs  
XX

KW cDNA vaccine; cytotoxic lymphocyte; peripheral blood.  
 XX Mus musculus.  
 OS US6514493-B1.  
 PN US6514493-B1.  
 XX US6514493-B1.  
 PD 04-FEB-2003.  
 XX 21-JUL-1997; 97US-00897843.  
 PF 21-JUL-1997; 97US-00897843.  
 PR 21-JUL-1997; 97US-00897843.  
 XX (UYPI-) UNIV PITTSBURGH.  
 PA Deleo AB, Loftus D, Appella E;  
 XX WPI; 2003-478760/45.  
 XX WPI; 2003-478760/45.  
 DR A new isolated cDNA molecule useful for the treatment of tumors comprises  
 XX a polynucleotide sequence encoding a protein fragment of murine  
 FT glycoprotein 110 containing 938 amino acids.  
 PT glycoprotein 110 containing 938 amino acids.  
 XX Example 7; Fig 1G; 27pp; English.  
 PS The invention relates to an isolated cDNA molecule (A) comprising a  
 XX polynucleotide sequence (not shown) encoding a protein fragment of murine  
 CC glycoprotein (gp) 110 appearing as ABU61901 (of 938 amino acids or their  
 CC conservative variants). Also included are a cDNA vaccine for inducing  
 CC resistance to tumors comprising (A) and an antigen presenting cell  
 CC transfected with (A). The cDNA is useful in the preparation of a vaccine  
 CC for inducing resistance to tumors in a patient (comprising transfecting  
 CC (A) into an antigen presenting cell (preferably a dendritic cell) and  
 CC administering the cell to the patient or distributing (A) on a particle  
 CC surface to form particulate polynucleotide and inoculating the patient  
 CC with the particulate polynucleotide). Variants of the gp peptide encoded  
 CC by the cDNA molecule are capable of inducing anti-gp 110 cytotoxic  
 CC lymphocytes in the peripheral blood of the normal individuals to induce  
 CC vaccination or in the treatment of tumours expressing gp 110 by causing  
 CC tumour rejection. The present sequence is a fragment of the mouse GP110  
 CC protein  
 XX Sequence 44 AA;  
 SQ  
 Query March 29.9%; Score 29; DB 6; Length 44;  
 Best Local Similarity 38.9%; Pred. No. 1.9e+03;  
 Matches 7; Conservative 4; Mismatches 5; Indels 2; Gaps 1;  
 QY 2 PNHLSKIAPKIVSQEPA 19  
 Db 25 PSHLN--LVFLSRAAA 40  
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 ID AAG59379 standard; protein; 45 AA.  
 XX  
 AC AAG59379;  
 XX  
 DT 18-OCT-2000 (first entry)  
 XX  
 DE Arabidopsis thaliana protein fragment SEQ ID NO: 76802.  
 XX  
 KW Protein identification; signal transduction pathway; metabolic pathway;  
 KW hybridisation assay; genetic mapping; gene expression control; promoter;  
 KW termination sequence.  
 XX Arabidopsis thaliana.  
 OS Arabidopsis thaliana.  
 XX EP1033405-A2.  
 PN 06-SEP-2000.  
 PD 25-FEB-2000; 2000EP-00301439.  
 XX  
 PF 25-FEB-1999;  
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 PR 01-JUL-1999;  
 PR 02-JUL-1999;  
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PR	15-JUL-1999;	99US-0144005P.
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PR	19-JUL-1999;	99US-0144333P.
PR	19-JUL-1999;	99US-0144334P.
PR	19-JUL-1999;	99US-0144335P.
PR	20-JUL-1999;	99US-0144352P.
PR	20-JUL-1999;	99US-0144632P.
PR	20-JUL-1999;	99US-0144884P.
PR	21-JUL-1999;	99US-0144814P.
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PR	23-JUL-1999;	99US-0145218P.
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PR	27-JUL-1999;	99US-0145918P.
PR	27-JUL-1999;	99US-0145919P.
PR	28-JUL-1999;	99US-0145951P.
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PR	29-SEP-1999;	99US-0156596P.
PR	04-OCT-1999;	99US-0157117P.
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PR	12-OCT-1999;	99US-0158369P.
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PR	13-OCT-1999;	99US-0159295P.
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PR	14-OCT-1999;	99US-0159331P.
PR	14-OCT-1999;	99US-0159637P.
PR	14-OCT-1999;	99US-0159638P.
PR	18-OCT-1999;	99US-0159584P.
PR	21-OCT-1999;	99US-0160741P.
PR	21-OCT-1999;	99US-0160767P.
PR	21-OCT-1999;	99US-0160768P.
PR	21-OCT-1999;	99US-0160770P.
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PR	21-OCT-1999;	99US-0160815P.
PR	22-OCT-1999;	99US-0160980P.
PR	22-OCT-1999;	99US-0160981P.
PR	22-OCT-1999;	99US-0160989P.
PR	25-OCT-1999;	99US-0161404P.
PR	25-OCT-1999;	99US-0161405P.
PR	25-OCT-1999;	99US-0161406P.
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PR	26-OCT-1999;	99US-0161361P.
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ID	ABBS0392 standard; protein; 46 AA.	
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AC	ABBS0392;	
XX		
DT	07-FEB-2002 (first entry)	
XX		
DE	Human secreted protein encoded by gene 92 SEQ ID NO:340.	
XX		
KW	Human; secreted protein; immunomodulatory; antisclerotic; anti-HIV;	
KW	dermatologic; immunosuppressive; antiinflammatory; immunostimulant;	
KW	cytostatic; cardiant; vascular; anti-angiogenic; ophthalmological;	
KW	neuroprotective; nootropic; anticonvulsant; antialzheimers; vulnery;	
KW	antiparkinsonian; antimicrobial; gene therapy; vaccine; immune disorder;	
KW	multiple sclerosis; systemic lupus erythematosus; HIV infection; cancer;	
KW	human immunodeficiency virus; hyperproliferative disorder; wound healing;	
KW	Chagaer's disease; cardiovascular disease; Scimitar syndrome; chemotaxis;	
KW	corneal graft neovascularisation; coronary arteriosclerosis; angioenic disorder;	
KW	neurological disorder; Huntington's chorea; Alzheimer's disease;	
XX	Parkinson's disease; infectious disease; chromosome 19.	
OS	Homo sapiens.	
XX		
PN	WC200162891-A2.	
PD	30-AUG-2001.	

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XX PF 21-FEB-2001; 2001WO-US005614.
XX PI Ni J, Ebner R, Lafleur DW, Moore PA, Olsen HS, Rosen CA;
XX PI Ruben SM, Soppet DR, Young PE, Shi Y, Florence KA, Wei Y;
XX PR Florence C, Hu J, Li Y, Kyaw H, Fischer CL, Ferrle AM, Fan P;
XX PR Feng P, Address GA, Dillon EV, Carter KC, Brewer LA, Yu G, Zeng Z;
XX PI Greene JM;
XX PA (HUMA-) HUMAN GENOME SCI INC.
XX NI WPI; 2001-625724/72.
XX DR N-PSDB; ABA83285.
XX NI Nucleic acids encoding 207 human secreted polypeptides, useful for
XX PT preventing, diagnosing and/or treating, e.g. cancers, Parkinson's disease
XX PT and diabetic retinopathy.
XX PS Claim 11; Page 1126; 1533pp; English.
XX CC ABB50301 to ABB51287 and ABA83194 to ABA83441 represent human secreted
XX CC proteins (I) and polynucleotide (II) sequences. (I) and (II) have various
XX CC activities based on the tissues and cells the genes are expressed in.
XX CC Example of these activities include: immunomodulatory; antisclerotic;
XX CC dermatological; immunosuppressive; antiinflammatory; immunostimulant;
XX CC anti-HIV; cytostatic; cardiac; anti-angiogenic; ophthalmological;
XX CC neuroprotective; nootropic; anticonvulsant; antialzheimers; vascular;
XX CC antiparkinsonian; antimicrobial; and vulnerary. (I) and (II) can be used
XX CC in gene therapy and vaccine production. (I) and (II) can be used in the
XX CC prevention, diagnosis and treatment of immune disorders (e.g. multiple
XX CC sclerosis, systemic lupus erythematosus and human immunodeficiency virus
XX CC (HIV) infections), hyperproliferative disorders (e.g. cancers and
XX CC Gaucher's disease), cardiovascular diseases (e.g. Schmittar syndrome,
XX CC Chaga's cardiomyopathy and coronary arteriosclerosis), angiogenic
XX CC disorders (e.g. corneal graft neovascularisation and diabetic
XX CC retinopathy), neurological disorders (e.g. Huntington's chorea,
XX CC Alzheimer's disease and Parkinson's disease), infectious diseases and/or
XX CC for promoting wound healing, regeneration and/or chemotaxis. ABA83185 to
XX CC ABA83193 and ABB50300 represent sequences used in the exemplification of
XX CC the present invention
XX SQ Sequence 46 AA;
    Query Match      29.9%; Score 29; DB 4; Length 46;
    Best Local Similarity 37.5%; Pred. No. 2e+03;
    Matches 6; Conservative 4; Mismatches 6; Indels 0; Gaps 0;
QY      3 NNLNSKIAFKIVSQEP 18
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XX AC ABO44649;
XX XX
XX DT 02-OCT-2003 (first entry)
XX DE Novel human secreted protein #92.
XX KW Human; gene therapy; autoimmune disorder; multiple sclerosis; cancer;
XX KW systemic lupus erythematosus; haematopoietic cell disorder; allergy;
XX KW agammaglobulinemia; ataxia telangiectasia; blood coagulation disorder;
XX KW afibrinogenemia; thrombocytopenia; graft-versus-host disease; arthritis;
XX KW inflammatory condition; ischaemia-reperfusion injury; infectious disease;
XX KW hyperproliferative disorder; purpura; viral infection; regeneration;
XX KW bacterial infection; ulcer; Alzheimer's disease.
XX XX

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OS Homo sapiens.
XX US2003065160-A1.
XX PD 03-APR-2003.
XX PF 07-DEC-2001; 2001US-00004860.
XX PR 06-JUN-1997; 97US-0048875P.
XX PR 06-JUN-1997; 97US-0048876P.
XX PR 06-JUN-1997; 97US-0048877P.
XX PR 06-JUN-1997; 97US-0048878P.
XX PR 06-JUN-1997; 97US-0048880P.
XX PR 06-JUN-1997; 97US-0048881P.
XX PR 06-JUN-1997; 97US-0048882P.
XX PR 06-JUN-1997; 97US-0048883P.
XX PR 06-JUN-1997; 97US-0048884P.
XX PR 06-JUN-1997; 97US-0048885P.
XX PR 06-JUN-1997; 97US-0048886P.
XX PR 06-JUN-1997; 97US-0048887P.
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XX PR 06-JUN-1997; 97US-0048889P.
XX PR 06-JUN-1997; 97US-0048890P.
XX PR 06-JUN-1997; 97US-0048901P.
XX PR 06-JUN-1997; 97US-0048915P.
XX PR 06-JUN-1997; 97US-0048916P.
XX PR 06-JUN-1997; 97US-0048917P.
XX PR 06-JUN-1997; 97US-0048949P.
XX PR 06-JUN-1997; 97US-0048962P.
XX PR 06-JUN-1997; 97US-0048963P.
XX PR 06-JUN-1997; 97US-0048964P.
XX PR 06-JUN-1997; 97US-0048970P.
XX PR 06-JUN-1997; 97US-0048971P.
XX PR 06-JUN-1997; 97US-0048972P.
XX PR 06-JUN-1997; 97US-0048974P.
XX PR 06-JUN-1997; 97US-0049019P.
XX PR 06-JUN-1997; 97US-0049020P.
XX PR 06-JUN-1997; 97US-0049373P.
XX PR 06-JUN-1997; 97US-0049374P.
XX PR 06-JUN-1997; 97US-0049375P.
XX PR 05-SEP-1997; 97US-0057584P.
XX PR 05-SEP-1997; 97US-0057627P.
XX PR 05-SEP-1997; 97US-0057628P.
XX PR 05-SEP-1997; 97US-0057629P.
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XX PR 05-SEP-1997; 97US-0057647P.
XX PR 05-SEP-1997; 97US-0057648P.
XX PR 05-SEP-1997; 97US-0057649P.
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XX PR 05-SEP-1997; 97US-0057651P.
XX PR 05-SEP-1997; 97US-0057654P.
XX PR 05-SEP-1997; 97US-0057661P.
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XX PR 05-SEP-1997; 97US-0057769P.

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PR 05-SEP-1997; 97US-0057770P.  
PR 05-SEP-1997; 97US-0057771P.  
PR 05-SEP-1997; 97US-0057774P.  
PR 05-SEP-1997; 97US-0057775P.  
PR 05-SEP-1997; 97US-0057776P.  
PR 05-SEP-1997; 97US-0057777P.  
PR 05-SEP-1997; 97US-0057778P.  
PR 18-DEC-1997; 97US-0070923P.  
PR 04-JUN-1998; 98WO-US011422.  
PR 15-JUL-1998; 98US-0092921P.  
PR 30-JUL-1998; 98US-0094657P.  
PR 04-DEC-1998; 98US-00205258.  
XX  
PA (HUMA-) HUMAN GENOME SCI INC.  
XX  
XX Young P, Greene JM, Ferrie AM, Ruben SM, Rosen CA, Hu J;  
PI Olsen HS, Ebner R, Brewer LA, Moore PA, Shi Y, Florence C;  
PI Florence K, Lafleur DW, Ni J, Fan P, Wei Y, Fischer CL, Soppet DR;  
PI Li Y, Zeng Z, Kyaw H, Yu G, Peng P, Dillon PJ, Endress GA;  
PI Carter KC;  
XX  
XX WPI; 2003-540804/51.  
DR N-PSDB; ACH04786.  
XX  
XX New isolated protein, useful for preparing a composition for diagnosing  
PT or treating cancer, inflammatory, immune or infectious diseases.  
XX  
XX Disclosure; SEQ ID NO 340; 172pp; English.  
XX  
XX The invention relates to an isolated HEMA80 protein. The protein is  
CC useful for preparing a composition for diagnosing or treating autoimmune  
CC disorders e.g. multiple sclerosis and systemic lupus erythematosus;  
CC haematopoietic cell disorders e.g. agammaglobulinaemia and ataxia  
CC telangiectasia; blood coagulation disorders e.g. afibrinogenaemia and  
CC thrombocytopenia; allergy; graft-versus-host disease; inflammatory  
CC conditions e.g. ischaemia-reperfusion injury and arthritis;  
CC hyperproliferative disorders e.g. cancer and purpura; infectious disease  
CC e.g. viral infection and bacterial infection. The polynucleotide or  
CC protein can be used to regenerate damaged tissue e.g. ulcers and  
CC Alzheimer's disease. The present sequence represents the amino acid  
CC sequence of a novel human secreted protein. Note: The sequence data for  
CC this patent did not form part of the printed specification but was  
CC obtained in electronic format directly from USPTO at  
CC seqdata.uspto.gov/sequence.html?DocID=20030065160  
XX  
XX Sequence 46 AA;  
SQ  
Query Match 29.9%; Score 29; DB 6; Length 46;  
Best Local Similarity 37.5%; Pred. No. 2e+03;  
Matches 6; Conservative 4; Mismatches 6; Indels 0; Gaps 0;  
QY 3 NHIHNSKIAFKIVSQEP 18  
DB 26 NLFTSQIKYIKYSEK 41  
RESULT 181  
ABO26129  
ID ABO26129 standard; protein; 46 AA.  
XX  
XX ABO26129;  
XX  
XX 10-SEP-2003 (first entry)  
XX  
XX Human protein from novel secreted protein gene 92.  
XX  
XX Human; secreted protein; precerebellin-like protein;  
KW neurodegenerative disorder; behavioural disorder; Alzheimer's disease;  
KW Parkinson's disease; Huntington's disease; schizophrenia; mania;  
KW dementia; paranoia; psychosis; autism; immune disorder; infection;  
KW inflammation; allergy; liver disorder; hepatoblastoma; jaundice;  
KW hepatitis; immunological disorder; AIDS; leukaemia; rheumatoid arthritis;  
KW sepsis; acne; psoriasis; cancer.

PR 05-SEP-1997; 97US-0057769P.  
 PR 05-SEP-1997; 97US-0057770P.  
 PR 05-SEP-1997; 97US-0057771P.  
 PR 05-SEP-1997; 97US-0057774P.  
 PR 05-SEP-1997; 97US-0057775P.  
 PR 05-SEP-1997; 97US-0057776P.  
 PR 05-SEP-1997; 97US-0057777P.  
 PR 05-SEP-1997; 97US-0057778P.  
 PR 18-DEC-1997; 97US-0070923P.  
 PR 04-JUN-1998; 98WO-US011422.  
 PR 15-JUL-1998; 98US-0092921P.  
 PR 30-JUL-1998; 98US-0094657P.  
 XX  
 PA (HUMA-) HUMAN GENOME SCI INC.  
 XX  
 PI Young P, Greene JM, Ferrie AM, Ruben SM, Rosen CA, Hu J;  
 PI Olsen HS, Ebner R, Brewer LA, Moore PA, Shi Y, Florence C;  
 PI Florence K, Lafleur DW, Ni J, Fan P, Wei Y, Fischer CL, Soppet DR;  
 PI Li Y, Zeng Z, Kyaw H, Yu G, Feng P, Dillon PJ, Endress GA;  
 PI Carter KC;  
 XX  
 DR WPI: 2003-511926/48.  
 DR N-PSDB; ACD44596.  
 XX  
 PT New precerebellin-like protein, useful for diagnosing or treating  
 PT neurodegenerative and behavioral disorders, immune disorders, liver  
 PT disorders, and cancer.  
 XX  
 PS Disclosure; SEQ ID NO 340; 156pp; English.  
 XX  
 CC The invention relates to an isolated protein comprising amino acid  
 CC residues 33-205 or 1-205 of a novel human secreted protein appearing as  
 CC ABO26252. The protein is encoded by one of 238 disclosed cDNA sequences  
 CC encoding 238 secreted proteins. ABO26252 is a precerebellin-like protein.  
 CC Also included are a composition comprising the protein and a carrier and  
 CC an isolated protein produced by expressing the protein cited above by a  
 CC cell, and recovering the protein. The proteins are useful for diagnosing  
 CC or treating neurodegenerative and behavioural disorders (e.g. Alzheimer's  
 CC disease, Parkinson's disease, Huntington's disease, schizophrenia, mania,  
 CC dementia, paranoia, psychoses or autism), immune disorders (e.g.  
 CC infection, inflammation, allergy), liver disorders (e.g. hepatoblastoma,  
 CC jaundice, hepatitis), immunological disorders (e.g. AIDS, leukaemia,  
 CC rheumatoid arthritis, sepsis, acne, psoriasis) and cancer. The present  
 CC sequence is one of the 238 disclosed novel secreted proteins. Note: The  
 CC sequence data for this patent did not form part of the printed  
 CC specification, but was obtained in electronic format directly from USPTO  
 CC at:- [seqdata.uspto.gov/sequence.html?DocID=6525174H1](http://seqdata.uspto.gov/sequence.html?DocID=6525174H1)  
 XX  
 SQ Sequence 46 AA;  
 Query Match 29.9%; Score 29; DB 7; Length 46;  
 Best Local Similarity 37.5%; Pred. No. 2e+03;  
 Matches 6; Conservative 4; Mismatches 6; Indels 0; Gaps 0;  
 QY 3 NHLNSKIAFKIVSQEP 18  
 Db 26 NLFTSQKIYKSEK 41  
 RESULT 182  
 AAB44907  
 ID AAB44907 standard; protein; 47 AA.  
 XX  
 AC AAB44907;  
 XX  
 XX 09-FEB-2001 (first entry)  
 DT  
 DE Human secreted protein encoded by gene 28 homologue.  
 XX  
 KW Human; secreted protein; cytostatic; antiarthritic; antiasthmatic;  
 KW immunosuppressive; antiarteriosclerotic; antiinflammatory; nootropic;  
 KW neuroprotective; antidiabetic; tranquiliser; vulnery; antibacterial;  
 KW antipsoriatic; antiarrhythmic; antirheumatic; cardiant; anti-HIV;

KW autoimmune disorder; allergic condition; cardiovascular disorder; cancer;  
 KW neurological disease; tissue repair.  
 XX  
 OS Homo sapiens.  
 PN WO200055176-A2.  
 XX  
 PD 21-SEP-2000.  
 XX  
 PF 09-MAR-2000; 2000WO-US006057.  
 XX  
 PR 12-MAR-1999; 99US-0124142P.  
 PR 11-JUN-1999; 99US-0138597P.  
 PR 03-DEC-1999; 99US-0168666P.  
 XX  
 XX (HUMA-) HUMAN GENOME SCI INC.  
 PA  
 PI Rosen CA, Ruben SM, Komatsoulis G;  
 PI  
 XX WPI: 2000-638176/61.  
 DR  
 XX  
 PT Novel 49 human secreted proteins useful for diagnosis, prevention and  
 PT treatment of disorders including neurological, cell proliferative,  
 PT cardiovascular, and autoimmune/inflammatory disorders and microbial  
 PT infections.  
 XX  
 PS Disclosure; Page 396; 405pp; English.  
 XX  
 CC This invention describes a novel isolated polypeptide (I) comprising an  
 CC amino acid sequence at least 95 % identical to a polypeptide sequence  
 CC selected from 49 polypeptides encoded by polynucleotide sequences  
 CC included in American Type Culture Collection (ATCC) deposit number  
 CC 203917, defined in the specification. The products of the invention have  
 CC cytostatic, antiarthritic, antiasthmatic, immunosuppressive, nootropic,  
 CC antiarteriosclerotic, antiinflammatory, neuroprotective, antidiabetic,  
 CC tranquiliser, vulnery, antibacterial, antipsoriatic, antiarrhythmic,  
 CC antirheumatic, cardiant and anti-HIV activity. (I) or a nucleic acid (II)  
 CC encoding (I) is useful for preventing, treating or ameliorating a medical  
 CC condition and for diagnosing a pathological condition or susceptibility  
 CC to the condition. (I) is useful for identifying a binding partner which  
 CC affects the activity of the polypeptide and for identifying an activity  
 CC in a biological sample. (I), (II) or an antibody (IV) specific to (I) is  
 CC also useful for treating or preventing a disease, disorder or condition  
 CC associated with aberrant expression of (I). Diseases treated or diagnosed  
 CC include immune disorders such as autoimmune diseases, blood protein  
 CC disorders, anemia, allergic reactions and conditions such as asthma,  
 CC organ rejection or graft-versus-host disease, inflammation, hyper  
 CC proliferative disorders, cardiovascular disorders such as arterioarterial  
 CC fistula, arrhythmias, arteriosclerosis, coronary thrombosis, organ  
 CC regeneration, cancer, neovascular glaucoma, diabetic retinopathy,  
 CC rheumatoid arthritis, psoriasis, diseases associated with increased  
 CC apoptosis that include acquired immunodeficiency syndrome (AIDS),  
 CC neurological diseases such as Parkinson's disease, viral, bacterial,  
 CC fungal or parasitic diseases. They are also used to repair, replace or  
 CC protect tissue damage by congenital defects, to treat trauma, in surgery,  
 CC including cosmetic plastic surgery, to treat fibrosis, reperfusion injury  
 CC or systemic cytokine damage, to stimulate chondrocyte growth, to prevent  
 CC skin aging due to sunburn, to change a mammal's mental state or physical  
 CC state by influencing biorhythms, cardiac rhythms, depression, memory,  
 CC stress and for accelerating wound healing. (I), (II) and/or their agonist  
 CC or antagonist are useful as food additives or preservatives to increase  
 CC or decrease storage capabilities, fat content, lipid, protein,  
 CC carbohydrate, vitamin, mineral or other nutritional components. (I) is  
 CC useful for screening therapeutic compounds. (II) is useful in forensic  
 CC biology for detecting DNA sequences and as diagnostic probes for  
 CC detecting the presence of specific mRNA in a particular cell type  
 XX  
 SQ Sequence 47 AA;  
 Query Match 29.9%; Score 29; DB 3; Length 47;  
 Best Local Similarity 71.4%; Pred. No. 2.1e+03;  
 Matches 5; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

QY 1 EPNHLS 7  
 Db 19 QENHLS 25

RESULT 183  
 ABG02562  
 ID ABG02562 standard; protein; 47 AA.  
 AC AC  
 XX AC  
 DT 13-FEB-2002 (first entry)  
 DE Novel human diagnostic protein #2553.  
 XX Human; chromosome mapping; gene mapping; gene therapy; forensic;  
 KW food supplement; medical imaging; diagnostic; genetic disorder.  
 XX Homo sapiens.  
 OS WO200175067-A2.  
 FN 11-OCT-2001.  
 PD 30-MAR-2001; 2001WO-US008631.  
 PF 31-MAR-2000; 2000US-00540217.  
 PR 23-AUG-2000; 2000US-00649167.  
 XX (HYSE-) HYSEQ INC.  
 PA Drmanac RT, Liu C, Tang YT;  
 PI WPI; 2001-639362/73.  
 DR N-PSDB; AAS66749.  
 XX New isolated polynucleotide and encoded polypeptides, useful in  
 PT diagnostics, forensics, gene mapping, identification of mutations  
 PT responsible for genetic disorders or other traits and to assess  
 PT biodiversity.  
 PS Claim 20; SEQ ID NO 32921; 103pp; English.  
 XX The invention relates to isolated polynucleotide (I) and polypeptide (II)  
 CC sequences. (I) is useful as hybridisation probes, polymerase chain  
 CC reaction (PCR) primers, oligomers, and for chromosome and gene mapping,  
 CC and in recombinant production of (II). The polynucleotides are also used  
 CC in diagnostics as expressed sequence tags for identifying expressed  
 CC genes. (I) is useful in gene therapy techniques to restore normal  
 CC activity of (II) or to treat disease states involving (II). (II) is  
 CC useful for generating antibodies against it, detecting or quantitating a  
 CC polypeptide in tissue, as molecular weight markers and as a food  
 CC supplement. (II) and its binding partners are useful in medical imaging  
 CC of sites expressing (II). (I) and (II) are useful for treating disorders  
 CC involving aberrant protein expression or biological activity. The  
 CC polypeptide and polynucleotide sequences have applications in  
 CC diagnostics, forensics, gene mapping, identification of mutations  
 CC responsible for genetic disorders or other traits to assess biodiversity  
 CC and to produce other types of data and products dependent on DNA and  
 CC amino acid sequences. ABG0010-ABG30377 represent novel human diagnostic  
 CC amino acid sequences of the invention. Note: The sequence data for this  
 CC patent did not appear in the printed specification, but was obtained in  
 CC electronic format directly from WIPO at  
 CC ftp.wipo.int/pub/published\_pct\_sequences  
 XX Sequence 47 AA;  
 SQ

Query Match 29.9%; Score 29; DB 4; Length 47;  
 Best Local Similarity 35.3%; Pred. No. 2.1e+03;  
 Matches 6; Conservative 3; Mismatches 8; Indels 0; Gaps 0;

QY 2 PNHLSKIAFKIVSQEP 18  
 Db 19 QENHLS 25

RESULT 184  
 AAM47156  
 ID AAM47156 standard; peptide; 47 AA.  
 AC AAM47156;  
 XX AC  
 DT 12-FEB-2002 (first entry)  
 DE Modular enzyme system related ACP-domain N-terminal peptide SRF2\_1.  
 XX Modular enzyme system; cyclic gene synthesis; repetitive coding sequence;  
 KW antibiotic; non-ribosomal peptide synthetase; NRPS; PKS;  
 XX polyketide synthase; actinomycin biosynthesis.  
 OS Bacillus subtilis.  
 XX WO200181564-A2.  
 FN 01-NOV-2001.  
 PD 25-APR-2001; 2001WO-DE001578.  
 PF 26-APR-2000; 2000DE-01021267.  
 PR (ACTI-) ACTINODRUG PHARM GMBH.  
 PA Schawecker F;  
 PI WPI; 2002-049276/06.  
 DR Preparing DNA encoding modular protein for e.g. producing new enzymes for  
 PT synthesis of polyketide antibiotics, comprises cyclic integration of  
 PT fragments into a vector.  
 XX Example 1; Fig 9; 83pp; German.  
 XX The present invention relates to the preparation of DNA, in a circular  
 CC vector, that encodes one or more segments of a modular polypeptide. DNA  
 CC or DNA libraries produced this way are used to produce modular  
 CC polypeptides, particularly enzymes, which can be used to act on  
 CC substrates to produce compounds for therapeutic testing. Enzymes of  
 CC particular interest are those involved in non-ribosomal peptide synthesis  
 CC or polyketide synthesis, and compounds for testing are particularly  
 CC macrocyclic antibiotics, including penicillins, vancomycins or  
 CC erythromycins, but may also be modular receptors. The present sequence is  
 CC a peptide used in the exemplification of the invention  
 SQ Sequence 47 AA;  
 SQ

Query Match 29.9%; Score 29; DB 5; Length 47;  
 Best Local Similarity 20.0%; Pred. No. 2.1e+03;  
 Matches 3; Conservative 7; Mismatches 5; Indels 0; Gaps 0;

QY 4 HLNSKIAFKIVSQEP 18  
 Db 20 HFKVPIKVLPEKP 34

RESULT 185  
 AAM47171  
 ID AAM47171 standard; peptide; 47 AA.  
 AC AAM47171;  
 XX AC  
 DT 29-AUG-2003 (revised)  
 DT 12-FEB-2002 (first entry)  
 XX Modular enzyme system related ACP-domain N-terminal peptide CVSA\_2.  
 DE Modular enzyme system; cyclic gene synthesis; repetitive coding sequence;  
 KW



KW antibiotic; non-ribosomal peptide synthetase; NRPS; PKS;  
 KW polyketide synthase; actinomycin biosynthesis.  
 XX  
 OS Tolypocladium inflatum.

PN WO200181564-A2.

XX  
 XX  
 PD 01-NOV-2001.

PF 25-APR-2001; 2001WO-DE001578.

PR 26-APR-2000; 2000DE-01021267.

XX (ACTI-) ACTINODRUG PHARM GMBH.

PI Schauwecker F;

DR WPI; 2002-049276/06.

XX Preparing DNA encoding modular protein for e.g. producing new enzymes for  
 PT synthesis of polyketide antibiotics, comprises cyclic integration of  
 PT fragments into a vector.

PS Example 1; Fig 9; 83pp; German.

XX The present invention relates to the preparation of DNA, in a circular  
 CC vector, that encodes one or more segments of a modular polypeptide. DNA  
 CC or DNA libraries produced this way are used to produce modular  
 CC polypeptides, particularly enzymes, which can be used to act on  
 CC substrates to produce compounds for therapeutic testing. Enzymes of  
 CC particular interest are those involved in non-ribosomal peptide synthesis  
 CC or polyketide synthesis, and compounds for testing are particularly  
 CC erythromycins, including penicillins, vancomycins or  
 CC macrolide antibiotics, but may also be modular receptors. The present sequence is  
 CC a peptide used in the exemplification of the invention. (Updated on 29-  
 CC AUG-2003 to standardise OS field)

XX Sequence 47 AA;

Query Match 29.9%; Score 29; DB 5; Length 47;  
 Best Local Similarity 42.9%; Pred. No. 2.1e+03;  
 Matches 6; Conservative 3; Mismatches 5; Indels 0; Gaps 0;

Qy 5 LNSKIAFKIVSQEP 18  
 ||:|||:  
 Db 21 LNAQIAVKDIFDRP 34

RESULT 186

AB054785  
 ID ABO54785 standard; protein; 47 AA.

XX  
 AC ABO54785;

XX  
 DT 29-JUL-2004 (first entry)

XX Human genome derived single exon protein #1019.

XX Human; gene expression; single exon probe; microarray;  
 KW alternative splicing event; genomic alteration.

XX Homo sapiens.

OS  
 XX US2003194704-A1.

XX  
 PD 16-OCT-2003.

XX 03-APR-2002; 2002US-00029386.

XX 03-APR-2002; 2002US-00029386.

XX (PENW/) PENN S G.

PA (RANK/) RANK D R.

PA (HANZ/) HANZEL D K.

XX Penn SG, Rank DR, Hanzel DK;

XX WPI; 2004-119264/12.

XX New human genome-derived single exon nucleic acid probes useful for human  
 PT gene expression analysis, for identifying or characterizing alternative  
 PT splicing events, for assessing genomic alterations or as tools for  
 PT surveying tissues.

XX Claim 45; SEQ ID NO 28419; 80pp; English.

XX The invention relates to a nucleic acid probe for measuring human gene  
 CC expression, comprising any of the 27,400 fully defined nucleotide  
 CC sequences in the specification, or their complements or fragments, and  
 CC encoding at least 8 amino acids of any of the 6888 amino acid sequences  
 CC fully defined in the specification. The probe is a single exon probe that  
 CC hybridises under high stringency conditions to a nucleic acid molecule  
 CC expressed in human cells or tissues. Also included are a spatially-  
 CC addressable set of single exon nucleic acid probes for measuring human  
 CC gene expression (comprising a plurality of single exon nucleic acid  
 CC probes cited above, where each of the plurality of probes is separately  
 CC and addressably isolatable or amplifiable from the plurality), a single  
 CC exon microarray for measuring human gene expression, a method of  
 CC measuring human gene expression, a vector comprising the single exon  
 CC probe cited above, an ORF-encoded peptide comprising at least 8  
 CC contiguous amino acids of any of the above-mentioned amino acid  
 CC sequences (optionally with conservative amino acid substitutions), an  
 CC isolated antibody that binds specifically to a peptide cited above,  
 CC methods of selling and/or licensing single exon probes or microarrays to  
 CC a customer desiring to measure gene expression, a method of providing  
 CC human gene expression data by subscription, and a computer-readable  
 CC storage medium which contains a database having a plurality of records  
 CC (each record including data on the expression of a single exon probe  
 CC cited above. The probe, methods and apparatus are useful in gene  
 CC expression analysis. The probes may be used as tools for surveying  
 CC tissues to detect the presence of expressed messages that contain their  
 CC specific exon, or in constructing genome-derived single exon microarrays.  
 CC In addition, the probes are used in identifying and characterising  
 CC alternative splicing events, in detecting and characterising gross  
 CC alterations in the genomic locus that includes their exon, in assessing  
 CC smaller genomic alterations, in priming the synthesis of nucleic acids,  
 CC or in expressing the ORF-encoded peptide. The present sequence is a human  
 CC single exon probe protein of the invention. Note: The sequence data for  
 CC this patent did not form part of the printed specification, but was  
 CC obtained in electronic format directly from USPTO at  
 CC seqdata.uspto.gov/sequence.html?DocID=20030194704

XX Sequence 47 AA;

Query Match 29.9%; Score 29; DB 8; Length 47;  
 Best Local Similarity 44.4%; Pred. No. 2.1e+03;  
 Matches 4; Conservative 4; Mismatches 1; Indels 0; Gaps 0;

Qy 2 PNHLNSKIA 10  
 |:|||:  
 Db 38 PSHLKSEVS 46

RESULT 187

AAB34265

ID AAB34265 standard; protein; 48 AA.

XX  
 AC AAB34265;

XX  
 DT 02-FEB-2001 (first entry)

XX Human secreted protein BLAST search protein SEQ ID NO: 111.

XX Cytostatic; immunosuppressive; nootropic; neuroprotective; antiviral;  
 KW antiallergic; hepatotropic; antidiabetic; antiinflammatory; antitumor;  
 KW vulnerary; anticonvulsant; antibacterial; antifungal; antiparasitic;

KW cardiant; gene therapy; cancer; immune disorder; cardiovascular disorder;  
 KW neurological disease; infection; human; secreted protein.

XX Homo sapiens.

XX WO2000055352-A2.

XX PD 21-SEP-2000.

XX PF 09-MAR-2000; 2000WO-US006044.

XX PR 12-MAR-1999; 99US-0124099P.

XX PR 03-DEC-1999; 99US-0168661P.

XX (HUMA-) HUMAN GENOME SCI INC.

XX PI Rosen CA, Ruben SM, Komatsoulis G;

XX WIPI; 2000-602124/57.

XX Novel human secreted proteins useful for diagnosis, prevention and  
 PT treatment of disorders including neurological, cell proliferative,  
 PT cardiovascular, autoimmune and inflammatory disorders and microbial  
 PT infections.

PS Disclosure; Page 366; 383pp; English.

XX The invention relates to the isolation of genes AAC59507-C59556 encoding  
 CC the human secreted proteins AAB34218-B34264. This sequence represents a  
 CC peptide fragment homologous to the protein encoded by the gene isolated  
 CC in the present invention. The sequence is a search result from a BLASTX  
 CC homology search. The genes and proteins are useful for preventing,  
 CC ameliorating or treating medical conditions, e.g. by protein or gene  
 CC therapy. The genes are isolated from a range of human tissues disclosed  
 CC in the specification. The nucleic acids, proteins, antibodies and  
 CC (ant)agonists are useful in the diagnosis, treatment and prevention of:  
 CC (a) cancer, e.g. breast and ovarian cancer, and other cancers of the  
 CC adrenal gland, bone, bone marrow, breast, gastrointestinal tract, liver,  
 CC lung, or urogenital; (b) immune disorders e.g. Addison's disease,  
 CC allergies, autoimmune haemolytic anaemia, autoimmune thyroiditis,  
 CC diabetes mellitus, Crohn's disease, multiple sclerosis, rheumatoid  
 CC arthritis and ulcerative colitis; (c) cardiovascular disorders such as  
 CC myocardial ischaemias; (d) wound healing; (e) neurological diseases e.g.  
 CC cerebral anoxia and epilepsy; and (f) infectious diseases such as viral,  
 CC bacterial, fungal and parasitic infections

XX SQ Sequence 48 AA;

Query Match 29.9%; Score 29; DB 3; Length 48;

Best Local Similarity 71.4%; Pred. No. 2.1e+03;

Matches 5; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

QY 1 EPNHLS 7

DB 17 QENHLS 23

RESULT 188

AAB58239

ID AAB58239 standard; protein; 48 AA.

XX AC AAB58239;

XX DT 14-MAR-2001 (first entry)

XX Lung cancer associated polypeptide sequence SEQ ID 577.

XX Human; lung cancer associated protein; neuroprotective; cytoskeletal;  
 KW cardioactive; immunomodulatory; muscular active; vulneryary;  
 KW gastrointestinal; nephrotropic; antiinfective; gynecological;  
 KW antibacterial; diagnosis; neural disorder; immune disorder; reproductive;  
 KW proliferative disorder; wound healing; infectious disease.

OS Homo sapiens.

XX PN WO2000055180-A2.

XX PD 21-SEP-2000.

XX PF 08-MAR-2000; 2000WO-US005918.

XX PR 12-MAR-1999; 99US-0124270P.

XX (HUMA-) HUMAN GENOME SCI INC.

XX (ROSE/) ROSEN C A.

XX PI Ruben SM;

XX WIPI; 2000-587514/55.

XX DR N-PSDB; AAF18115.

XX Lung cancer associated gene sequences, referred to as lung cancer  
 PT antigens, useful for treatment, prevention, and diagnosis of disorders  
 PT such as lung cancer.

PS Claim 11; Page 1072; 1425pp; English.

XX Polynucleotide sequences AAF17982 - AAF18424 encode human lung cancer  
 CC associated proteins represented in AAB58106 - AAB58548. Lung cancer  
 CC associated proteins and polynucleotide sequences, their agonists, and  
 CC antagonists may have neuroprotective; cytoskeletal; cardioactive;  
 CC immunomodulatory; muscular active general; vulneryary; gastrointestinal  
 CC general; nephrotropic; antiinfective; gynecological; or antibacterial  
 CC activity. The invention also includes antibodies specific for the protein  
 CC or polynucleotide sequences. The lung cancer associated polynucleotide  
 CC sequences may be used for detection of lung cancer, chromosome  
 CC identification, as chromosome markers, and for numerous other diagnostic  
 CC or research purposes. The proteins may be used to treat disorders such as  
 CC neural, immune, muscular, reproductive, gastrointestinal, pulmonary,  
 CC cardiovascular, renal, and proliferative disorders. The proteins may also  
 CC be used in the treatment of wounds and infectious diseases.

CC Polynucleotide sequences AAF18425 - AAF18433 and peptide AAB58549 are  
 CC used in the course of the invention for the identification and  
 CC characterisation of the polynucleotide and protein sequences

XX SQ Sequence 48 AA;

Query Match 29.9%; Score 29; DB 3; Length 48;

Best Local Similarity 42.9%; Pred. No. 2.1e+03;

Matches 6; Conservative 3; Mismatches 5; Indels 0; Gaps 0;

QY 5 LNSKJAFKIVSQEP 18

DB 6 JHSSLAKKLKNEP 19

RESULT 189

AAG73655

ID AAG73655 standard; protein; 48 AA.

XX AC AAG73655;

XX DT 03-SEP-2001 (first entry)

XX Human colon cancer antigen protein SEQ ID NO:4419.

XX Human; colon cancer; colon cancer antigen; diagnosis; detection;  
 KW colorectal carcinoma.

XX OS Homo sapiens.

XX WO200122920-A2.

XX PD 05-APR-2001.

XX PF 28-SEP-2000; 2000WO-US026524.

XX 29-SEP-1999; 99US-0157137P.  
 PR 03-NOV-1999; 99US-0163280P.  
 XX (HUMA-) HUMAN GENOME SCI INC.  
 XX Ruben SM, Barash SC, Birse CE, Rosen CA;  
 PI WPI; 2001-235357/24.  
 DR N-PSDB; AAH33086.  
 XX Nucleic acids encoding 4277 human colon cancer-associated polypeptides,  
 PT useful for preventing, diagnosing and/or treating colorectal cancers.  
 XX Claim 11; Page 6239; 9803pp; English.  
 XX AAH32943 to AAH37195 and AAG73514 to AAG77788 represent human colon  
 CC cancer-associated nucleic acid molecules (N) and proteins (P), where the  
 CC proteins are collectively known as colon cancer antigens. The colon  
 CC cancer antigens have cytostatic activity and can be used in gene therapy  
 CC and vaccine production. N and P may be used in the prevention, diagnosis  
 CC and treatment of diseases associated with inappropriate P expression. For  
 CC example, N and P may be used to treat disorders associated with decreased  
 CC expression by rectifying mutations or deletions in a patient's genome  
 CC that affect the activity of P by expressing inactive proteins or to  
 CC supplement the patients own production of P. Additionally, N may be used  
 CC to produce the colon cancer-associated Ps, by inserting the nucleic acids  
 CC into a host cell and culturing the cell to express the proteins. N and P  
 CC can be used in the prevention, diagnosis and treatment of colorectal  
 CC carcinomas and cancers. AAH37196 to AAH37204 and AAB77789 represent  
 CC sequences used in the exemplification of the present invention. N.B.  
 CC Pages 666 to 682 and page 7053 of the sequence listing were missing at  
 CC time of publication, meaning no sequences are present for SEQ ID NO:1027  
 CC to 1052, 7921 and 7922  
 XX Sequence 48 AA;  
 SQ Query Match 29.9%; Score 29; DB 4; Length 48;  
 Best Local Similarity 41.7%; Pred. No. 2.1e+03;  
 Matches 5; Conservative 3; Mismatches 4; Indels 0; Gaps 0;  
 Qy 3 NHLNLSKIAFKIV 14  
 Db ||| | : | : |  
 10 NHNCYILYFII 21  
 RESULT 190  
 ABB01426  
 ID ABB01426 standard; protein; 48 AA.  
 XX AC ABB01426;  
 XX DT 13-FEB-2002 (first entry)  
 XX Novel human diagnostic protein #1417.  
 XX Human; chromosome mapping; gene mapping; gene therapy; forensic;  
 KW food supplement; medical imaging; diagnostic; genetic disorder.  
 XX Homo sapiens.  
 OS WO200175067-A2.  
 XX PN 11-OCT-2001.  
 XX PF 30-MAR-2001; 2001WO-US008631.  
 XX PR 31-MAR-2000; 2000US-00540217.  
 XX PR 23-AUG-2000; 2000US-00649167.  
 XX PA (HYSE-) HYSEQ INC.  
 XX Drmanac RT, Liu C, Tang YT;

XX WPI; 2001-639362/73.  
 DR N-PSDB; AAS65613.  
 XX New isolated polynucleotide and encoded polypeptides, useful in  
 PT diagnostics, forensics, gene mapping, identification of mutations  
 PT responsible for genetic disorders or other traits and to assess  
 PT biodiversity.  
 XX Claim 20; SEQ ID NO 31785; 103pp; English.  
 XX The invention relates to isolated polynucleotide (I) and polypeptide (II)  
 CC sequences. (I) is useful as hybridisation probes, polymerase chain  
 CC reaction (PCR) primers, oligomers, and for chromosome and gene mapping,  
 CC and in recombinant production of (II). The polynucleotides are also used  
 CC in diagnostics as expressed sequence tags for identifying expressed  
 CC genes. (I) is useful in gene therapy techniques to restore normal  
 CC activity of (II) or to treat disease states involving (II). (II) is  
 CC useful for generating antibodies against it, detecting or quantitating a  
 CC polypeptide in tissue, as molecular weight markers and as a food  
 CC supplement. (II) and its binding partners are useful in medical imaging  
 CC of sites expressing (II). (I) and (II) are useful for treating disorders  
 CC involving aberrant protein expression or biological activities. The  
 CC polypeptide and polynucleotide sequences have applications in  
 CC diagnostics, forensics, gene mapping, identification of mutations  
 CC responsible for genetic disorders or other traits to assess biodiversity  
 CC and to produce other types of data and products dependent on DNA and  
 CC amino acid sequences. ABB00010-ABG30377 represent novel human diagnostic  
 CC amino acid sequences of the invention. Note: The sequence data for this  
 CC patent did not appear in the printed specification, but was obtained in  
 CC electronic format directly from WIPO at  
 CC ftp.wipo.int/pub/published\_pct\_sequences  
 XX Sequence 48 AA;  
 SQ Query Match 29.9%; Score 29; DB 4; Length 48;  
 Best Local Similarity 55.6%; Pred. No. 2.1e+03;  
 Matches 5; Conservative 3; Mismatches 1; Indels 0; Gaps 0;  
 Qy 5 LNSKIAFKI 13  
 Db ||| | : | : |  
 36 VNSMVAYKI 44  
 RESULT 191  
 AAB32117  
 ID AAB32117 standard; protein; 49 AA.  
 XX AC AAB32117;  
 XX DT 14-FEB-2001 (first entry)  
 XX Human secreted protein BLAST search protein SEQ ID NO: 175.  
 DE Cytostatic; immunosuppressive; nootropic; neuroprotective; antiviral;  
 KW antiallergic; hepatotropic; antidiabetic; antiinflammatory; antiulcer;  
 KW vulnery; anticonvulsant; antibacterial; antifungal; antiparasitic;  
 KW cardiant; gene therapy; cancer; immune disorder; cardiovascular disorder;  
 KW neurological disease; infection; human; secreted protein.  
 XX Homo sapiens.  
 OS WO200058350-A1.  
 XX PN 05-OCT-2000.  
 XX PD 22-MAR-2000; 2000WO-US007483.  
 XX PF 26-MAR-1999; 99US-0126596P.  
 XX PR 22-DEC-1999; 99US-0171552P.  
 XX (HUMA-) HUMAN GENOME SCI INC.  
 XX

PI Rosen CA, Ruben SM, Komatsoulis G;  
 XX WPI; 2000-602357/57.  
 XX Nucleic acid molecules encoding human secreted proteins, used in  
 PT preventing, treating or ameliorating a disorder, e.g. Alzheimer's and  
 PT Parkinson's diseases and cancers.  
 XX Disclosure; Page 420; 423pp; English.  
 XX The invention relates to the isolation of genes AAC66410-C66458 encoding  
 CC the human secreted proteins AAB32002-B32050. This sequence represents a  
 CC peptide fragment homologous to the protein encoded by the gene given in  
 CC the descriptor line. The sequence is a search result from a BLASTX  
 CC homology search. The genes and proteins are useful for preventing,  
 CC ameliorating or treating medical conditions, e.g. by protein or gene  
 CC therapy. The genes are isolated from a range of human tissues disclosed  
 CC in the specification. The nucleic acids, proteins, antibodies and  
 CC (ant)agonists are useful in the diagnosis, treatment and prevention of:  
 CC (a) cancer, e.g. breast and ovarian cancer, and other cancers of the  
 CC adrenal gland, bone, bone marrow, breast, gastrointestinal tract, liver,  
 CC lung, or urogenital; (b) immune disorders e.g. Addison's disease,  
 CC allergies, autoimmune haemolytic anaemia, autoimmune thyroiditis,  
 CC diabetes mellitus, Crohn's disease, multiple sclerosis, rheumatoid  
 CC arthritis and ulcerative colitis; (c) cardiovascular disorders such as  
 CC myocardial ischaemias; (d) wound healing; (e) neurological diseases e.g.  
 CC cerebral anoxia and epilepsy; and (f) infectious diseases such as viral,  
 CC bacterial, fungal and parasitic infections  
 XX Sequence 49 AA;  
 SQ Query Match 29.9%; Score 29; DB 3; Length 49;  
 Best Local Similarity 71.4%; Pred. No. 2.2e+03;  
 Matches 5; Conservative 1; Mismatches 1; Indels 0; Gaps 0;  
 QY 1 EPNHLS 7  
 Db 19 QENHLS 25  
 RESULT 192  
 AAB45192  
 ID AAB45192 standard; protein; 50 AA.  
 XX AAB45192;  
 AC AAB45192;  
 XX 12-FEB-2001 (first entry)  
 DT Gene 26 human secreted protein homologous amino acid sequence #133.  
 DE Human; secreted protein; immunosuppressive; antiarthritic; antirheumatic;  
 KW antiproliferative; cytostatic; cardiant; vasotropic; cerebroprotective;  
 KW nootropic; neuroprotective; antibacterial; virucide; fungicide; cancer;  
 KW opthalmological; autoimmune disease; hyperproliferative disorder;  
 KW cardiovascular disorder; cerebrovascular disorder; wound healing;  
 KW nervous system disorder; aging; chemotaxis.  
 XX Homo sapiens.  
 OS Homo sapiens.  
 XX WO200058467-A1.  
 PN 05-OCT-2000.  
 PD 22-MAR-2000; 2000WO-US007505.  
 PF 26-MAR-1999; 99US-0126502P.  
 PR 17-DEC-1999; 99US-0172410P.  
 XX (HUMA-) HUMAN GENOME SCI INC.  
 PA Rosen CA, Ruben SM, Komatsoulis G;  
 XX WPI; 2000-611712/58.  
 DR

XX Nucleic acid molecules encoding human secreted proteins, used in  
 PT preventing, treating or ameliorating a disorder, e.g. Alzheimer's and  
 PT Parkinson's diseases and cancers.  
 XX Disclosure; Page 40; 440pp; English.  
 XX Polynucleotide sequences AAC80531-C80580 represent cDNA encoding human  
 CC secreted proteins AAB45120-B45169. Sequences AAB45170-B45225 represent  
 CC alternative polypeptides encoded by the genes, and amino acid sequences  
 CC to which they are homologous. The genes and proteins have activities  
 CC dependent on the tissues and cells in which they are expressed. Examples  
 CC of their activities include immunosuppressive; antiarthritic;  
 CC antirheumatic; antiproliferative; cytostatic; cardiant; vasotropic;  
 CC cerebroprotective; nootropic; neuroprotective; antibacterial; virucide;  
 CC fungicide; and opthalmological. The secreted proteins, polynucleotides,  
 CC antagonists and agonists may be useful in treating, preventing and/or  
 CC diagnosing diseases and disorders such as autoimmune diseases e.g.  
 CC rheumatoid arthritis, hyperproliferative disorders e.g. neoplasms of the  
 CC breast or liver, cardiovascular disorders e.g. cardiac arrest,  
 CC cerebrovascular disorders e.g. cerebral ischaemia, angiogenesis, nervous  
 CC system disorders e.g. Alzheimer's disease, infections caused by bacteria,  
 CC viruses and fungi and ocular disorders e.g. corneal infection. The  
 CC polypeptides can also be used to aid wound healing and epithelial cell  
 CC proliferation, to prevent skin aging due to sunburn, to maintain organs  
 CC before transplantation, for supporting cell culture of primary tissues,  
 CC to regenerate tissues and in chemotaxis. The polypeptides can also be  
 CC used as a food additive or preservative to increase or decrease storage  
 CC capabilities. AAC80522-C80530 and AAB45119 represent sequences used in  
 CC the isolation and characterisation of the genes and proteins of the  
 CC invention  
 XX Sequence 50 AA;  
 SQ Query Match 29.9%; Score 29; DB 3; Length 50;  
 Best Local Similarity 71.4%; Pred. No. 2.2e+03;  
 Matches 5; Conservative 1; Mismatches 1; Indels 0; Gaps 0;  
 QY 1 EPNHLS 7  
 Db 15 QENHLS 21  
 RESULT 193  
 AAB29913  
 ID AAB29913 standard; protein; 50 AA.  
 XX AAB29913;  
 AC AAB29913;  
 XX 09-FEB-2001 (first entry)  
 DT Human secreted protein BLAST search protein SEQ ID NO: 171.  
 DE Cytostatic; immunosuppressive; nootropic; neuroprotective; antiviral;  
 KW antiallergic; hepatotropic; antidiabetic; antiinflammatory; antitumor;  
 KW vulnary; anticonvulsant; antibacterial; antifungal; antiparasitic;  
 KW cardiant; gene therapy; cancer; immune disorder; cardiovascular disorder;  
 KW neurological disease; infection; human; secreted protein.  
 XX Homo sapiens.  
 OS Homo sapiens.  
 XX WO200061779-A1.  
 PN 19-OCT-2000.  
 PD 06-APR-2000; 2000WO-US009068.  
 PF 09-APR-1999; 99US-0128699P.  
 PR 20-JAN-2000; 2000US-0177050P.  
 XX (HUMA-) HUMAN GENOME SCI INC.  
 PA Rosen CA, Ruben SM, Komatsoulis G;  
 XX WPI; 2000-611712/58.  
 DR

XX WPI; 2000-647424/52.  
 XX Isolated nucleic acid molecule encoding a human secreted protein is used  
 PT in preventing, treating or ameliorating a medical condition.  
 XX  
 XX Disclosure; Page 489-490; 495pp; English.  
 PS  
 CC The invention relates to the isolation of genes AAC63410-C63458 encoding  
 CC the human secreted proteins AB29802-B29850. This sequence represents a  
 CC peptide fragment homologous to the protein encoded by the gene given in  
 CC the descriptor line. The sequence is a search result from a BLASTX  
 CC homology search. The genes and proteins are useful for preventing,  
 CC ameliorating or treating medical conditions, e.g. by protein or gene  
 CC therapy. The genes are isolated from a range of human tissues disclosed  
 CC in the specification. The nucleic acids, proteins, antibodies and  
 CC (ant)agonists are useful in the diagnosis, treatment and prevention of:  
 CC (a) cancer, e.g. breast and ovarian cancer, and other cancers of the  
 CC adrenal gland, bone, bone marrow, breast, gastrointestinal tract, liver,  
 CC lung, or urogenital; (b) immune disorders e.g. Addison's disease,  
 CC allergies, autoimmune haemolytic anaemia, autoimmune thyroiditis,  
 CC diabetes mellitus, Crohn's disease, multiple sclerosis, rheumatoid  
 CC arthritis and ulcerative colitis; (c) cardiovascular disorders such as  
 CC myocardial ischaemias; (d) wound healing; (e) neurological diseases e.g.  
 CC cerebral anoxia and epilepsy; and (f) infectious diseases such as viral,  
 CC bacterial, fungal and parasitic infections  
 XX  
 SQ Sequence 50 AA;  
 Query Match 29.9%; Score 29; DB 3; Length 50;  
 Best Local Similarity 71.4%; Pred. No. 2.2e+03;  
 Matches 5; Conservative 1; Mismatches 1; Indels 0; Gaps 0;  
 QY 1 EPNHLNS 7  
 Db 36 QENHLNS 42  
 RESULT 194  
 ABP29303  
 ID ABP29303 standard; protein; 50 AA.  
 XX AC ABP29303;  
 XX DT 02-JUL-2002 (first entry)  
 XX DE Streptococcus polypeptide SEQ ID NO 7782.  
 XX Streptococcus; GAS; GBS; group B streptococcus; Streptococcus agalactiae;  
 XX group A streptococcus; Streptococcus pyogenes; antibacterial;  
 XX antiinflammatory; infection; vaccine; meningitis; gene therapy.  
 XX Streptococcus pyogenes.  
 XX WO200234771-A2.  
 XX 02-MAY-2002.  
 XX 29-OCT-2001; 2001WO-GB004789.  
 XX 27-OCT-2000; 2000GB-00026333.  
 XX 24-NOV-2000; 2000GB-00028727.  
 XX 07-MAR-2001; 2001GB-00005640.  
 XX (CHIR-) CHIRON SPA.  
 XX (GENO-) INST GENOMIC RES.  
 XX Telford J, Masignani V, Margarit Y RosI, Grandi G, Fraser C;  
 XX Tettelin H;  
 XX WPI; 2002-352536/38.  
 XX N-PSDB; ABN69934.

PT New Streptococcus protein for the treatment or prevention of infection or  
 PT disease caused by Streptococcus bacteria, such as meningitis, and for  
 XX detecting a compound that binds to the protein.  
 PS Claim 1; Page 3913; 4525pp; English.  
 XX  
 CC The invention relates to a protein (ABP25413-ABP30895) from group B  
 CC Streptococcus/GBS (Streptococcus agalactiae) or group A streptococcus/GAS  
 CC (Streptococcus pyogenes), comprising one of 5483 sequences (S1), given in  
 CC the specification. The proteins have antibacterial and antiinflammatory  
 CC activity. (I), nucleic acids encoding (I), ABN66044-ABN71526 and  
 CC antibodies that bind (I) are used in the manufacture of medicaments for  
 CC the treatment or prevention of infection or disease caused by  
 CC Streptococcus bacteria, particularly S. agalactiae and S. pyogenes.  
 CC Nucleic acids encoding (I) are used to detect Streptococcus in a  
 CC biological sample. (I) is used to determine whether a compound binds to  
 CC (I). A composition comprising (I) or a nucleic acid encoding (I), may be  
 CC used as a vaccine or diagnostic composition. The disease caused by  
 CC Streptococcus that is prevented or treated may be meningitis. Nucleic  
 CC acid encoding (I) may be used to recombinantly produce (I) and may be  
 CC used in gene therapy. Antibodies to (I) are used for affinity  
 CC chromatography, immunoassays, and distinguishing/identifying  
 CC Streptococcus proteins  
 XX  
 SQ Sequence 50 AA;  
 Query Match 29.9%; Score 29; DB 5; Length 50;  
 Best Local Similarity 50.0%; Pred. No. 2.2e+03;  
 Matches 6; Conservative 1; Mismatches 5; Indels 0; Gaps 0;  
 QY 4 HLNSKIAPKIVS 15  
 Db 25 HSNNSICINIVS 36  
 RESULT 195  
 ADD90406  
 ID ADD90406 standard; protein; 50 AA.  
 XX AC ADD90406;  
 XX DT 29-JAN-2004 (first entry)  
 XX DE Novel human secreted protein seq id 26 protein feature seq id 221.  
 XX gene therapy; cytostatic; cancer; human; secreted protein.  
 XX Homo sapiens.  
 XX US2003199683-A1.  
 XX 23-OCT-2003.  
 XX 30-MAR-2001; 2001US-00820649.  
 XX 30-JUL-1997; 97US-0054209P.  
 XX 30-JUL-1997; 97US-0054211P.  
 XX 30-JUL-1997; 97US-0054212P.  
 XX 30-JUL-1997; 97US-0054213P.  
 XX 30-JUL-1997; 97US-0054214P.  
 XX 30-JUL-1997; 97US-0054215P.  
 XX 30-JUL-1997; 97US-0054217P.  
 XX 30-JUL-1997; 97US-0054218P.  
 XX 30-JUL-1997; 97US-0054234P.  
 XX 30-JUL-1997; 97US-0054236P.  
 XX 18-AUG-1997; 97US-0055968P.  
 XX 18-AUG-1997; 97US-0055969P.  
 XX 18-AUG-1997; 97US-0055972P.  
 XX 19-AUG-1997; 97US-0056534P.  
 XX 19-AUG-1997; 97US-0056543P.  
 XX 19-AUG-1997; 97US-0056554P.  
 XX 19-AUG-1997; 97US-0056561P.  
 XX 19-AUG-1997; 97US-0056727P.

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PR 19-AUG-1997; 97US-0056729P.
PR 19-AUG-1997; 97US-0056730P.
PR 29-JUL-1998; 98WO-US015949.
PR 26-JAN-1999; 99US-00236557.
PR 21-SEP-2000; 2000US-00666987.
XX (RUBE/) RUBEN S M.
PA (FENG/) FENG P.
PA (LAFLE/) LAFLEUR D W.
PA (MOOR/) MOORE P A.
PA (SHI/) SHI Y.
PA (KYAW/) KYAW H.
PA (LIY/) LI Y.
PA (ZENG/) ZENG Z.
PA (CART/) CARTER K C.
PA (ENDR/) ENDRESS G A.
PA (WEI/) WEI Y.
PA (FANP/) FAN P.
PA (ROSE/) ROSEN C A.
XX
XX Ruben SM, Feng P, Lafleur DW, Moore PA, Shi Y, Kyaw H, Li Y;
PI Zeng Z, Carter KC, Endress GA, Wei Y, Fan P, Rosen CA;
XX WPI; 2003-852813/79.
XX
XX New nucleic acid molecule, useful for preparing a medicament for
PT preventing, treating or ameliorating a medical condition e.g., cancer.
XX
XX Disclosure; SEQ ID NO 221; 213pp; English.
XX
XX The invention describes novel isolated human nucleic acids. The nucleic
CC acid is useful for preparing a medicament for preventing, treating or
CC ameliorating a medical condition e.g., cancer, and in gene therapy. This
CC is the amino acid sequence of polypeptide feature of a novel human
CC secreted protein of the invention.
XX
XX Sequence 50 AA;
SQ
Query Match 29.9%; Score 29; DB 7; Length 50;
Best Local Similarity 38.5%; Pred. No. 2.2e+03;
Matches 5; Conservative 3; Mismatches 5; Indels 0; Gaps 0;
QY 2 PNHLNSKIAFKIV 14
Db 30 PDHVNMSLVKII 42
RESULT 196
ADG90225
ID ADG90225 standard; protein; 50 AA.
XX
XX AC ADG90225;
XX
XX DT 11-MAR-2004 (first entry)
XX
XX DE Human secreted protein gene 16 extra polypeptide #2.
XX
XX Secreted protein; gene therapy; neural disorder; immune system disorders;
KW muscular disorder; reproductive disorder; gastrointestinal disorder;
KW pulmonary disorder; cardiovascular disorder; renal disorder;
KW proliferative disorder; cancer; systemic lupus erythematosus;
KW rheumatoid arthritis; multiple sclerosis; thyroiditis; anaemia;
KW Grave's disease; diabetes; hepatitis; asthma; allergy; nephritis;
KW Parkinson's disease; Alzheimer's disease; atherosclerosis;
KW myocardial infarction; AIDS; infection; human.
XX
XX OS Homo sapiens.
XX
XX PN US2003166541-A1.
XX
XX PD 04-SEP-2003.
XX
XX PF 04-JUN-2002; 2002US-00160162.
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XX 30-JUL-1997; 97US-0054209P.
PR 30-JUL-1997; 97US-0054211P.
PR 30-JUL-1997; 97US-0054212P.
PR 30-JUL-1997; 97US-0054213P.
PR 30-JUL-1997; 97US-0054214P.
PR 30-JUL-1997; 97US-0054215P.
PR 30-JUL-1997; 97US-0054217P.
PR 30-JUL-1997; 97US-0054218P.
PR 30-JUL-1997; 97US-0054234P.
PR 30-JUL-1997; 97US-0054236P.
PR 18-AUG-1997; 97US-0055968P.
PR 18-AUG-1997; 97US-0055969P.
PR 19-AUG-1997; 97US-0056534P.
PR 19-AUG-1997; 97US-0056543P.
PR 19-AUG-1997; 97US-0056554P.
PR 19-AUG-1997; 97US-0056561P.
PR 19-AUG-1997; 97US-0056727P.
PR 19-AUG-1997; 97US-0056729P.
PR 19-AUG-1997; 97US-0056730P.
PR 29-JUL-1998; 98WO-US015949.
PR 26-JAN-1999; 99US-00236557.
PR 05-JUN-2001; 2001US-0295558P.
XX (HUMA-) HUMAN GENOME SCI INC.
XX
XX Ruben SM, Feng P, Lafleur DW, Moore PA, Shi Y, Kyaw H, Li Y;
PI Zeng Z, Carter KC, Endress GA, Wei Y, Fan P, Rosen CA;
XX WPI; 2003-874923/81.
XX
XX Nucleic acids encoding 83 secreted polypeptides, useful for preventing,
PT diagnosing and treating disorders related to their aberrant expression
PT and activity.
XX
XX Disclosure; SEQ ID NO 221; 308pp; English.
XX
XX The invention relates to an isolated nucleic acid molecule encoding a
CC secreted protein that is at least 95% identical to a polynucleotide
CC fragment of any of the nucleotide sequences listed in table 1A of the
CC specification, which is hybridisable to the nucleotide sequences, a
CC polynucleotide encoding a polypeptide (or a polypeptide fragment, domain
CC or epitope of any of the amino acid sequences) listed in table 1A of the
CC specification, a polynucleotide which is an (allelic) variant of the
CC nucleotide sequences listed in the specification, a polynucleotide which
CC encodes a species homologue of the above amino acid sequences, a
CC polynucleotide capable of hybridising under stringent conditions to any
CC of the above polynucleotides, where the polynucleotide does not hybridise
CC under stringent conditions to a nucleic acid molecule having a nucleotide
CC sequence of only A or T residues. Also included are a recombinant vector
CC comprising the above nucleic acid molecule, making a recombinant host
CC cell comprising the above nucleic acid molecule, an isolated polypeptide
CC comprising a sequence that is at least 95% identical to the polypeptide
CC (or its fragment, domain, epitope, secreted form, (allelic) variant or
CC homologue) encoded by the above nucleic acid molecule, an isolated
CC antibody that binds specifically to the above polypeptide, a recombinant
CC host cell produced by the above method and that expresses the above
CC polypeptide, making an isolated polypeptide, preventing, treating or
CC ameliorating a medical condition, diagnosing a pathological condition or
CC a susceptibility to a pathological condition in a subject, identifying a
CC binding partner to the above polypeptide, and the gene corresponding to the
CC cDNA sequence given in the specification, and identifying an activity in
CC a biological assay. The nucleic acid molecule and polypeptide are useful
CC in diagnosing, preventing, prognosing or treating diseases or disorders
CC associated with aberrant expression and/or activity of the above
CC polypeptide, such as neural disorders, immune system disorders, muscular
CC disorders, reproductive disorders, gastrointestinal disorders, pulmonary
CC disorders, cardiovascular disorders, renal disorders, proliferative
CC disorders and/or cancers. In particular, these diseases are systemic
CC lupus erythematosus, rheumatoid arthritis, multiple sclerosis,
CC thyroiditis, anaemia, Grave's disease, diabetes, hepatitis, asthma,
CC allergies, nephritis, Parkinson's disease, Alzheimer's disease,
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CC atherosclerosis, myocardial infarction, AIDS and infections. The methods  
 CC may be used for identifying agonists and antagonists of the  
 CC polynucleotide and polypeptide. The present sequence is a protein from  
 CC one of the 83 disclosed secreted protein genes.

XX Sequence 50 AA;

Query Match 29.9%; Score 29; DB 7; Length 50;  
 Best Local Similarity 38.5%; Pred. No. 2.2e+03;  
 Matches 5; Conservative 3; Mismatches 5; Indels 0; Gaps 0;

Qy 2 PNHLSKIAFKIV 14  
 :|:|:  
 Db 30 PDHVNMSLVKII 42

RESULT 197  
 AAM33917  
 ID AAM33917 standard; protein; 30 AA.

XX AC AAM33917;

DT 17-OCT-2001 (first entry)

XX Peptide #7954 encoded by probe for measuring placental gene expression.

DE Probe; microarray; human; placenta; antenatal diagnosis;  
 XX genetic disorder.

KW Homo sapiens.

OS WO200157272-A2.

XX PN 09-AUG-2001.

XX PF 30-JAN-2001; 2001WO-US000663.

XX PR 04-FEB-2000; 2000US-0180312P.

XX PR 26-MAY-2000; 2000US-0207456P.

XX PR 30-JUN-2000; 2000US-00608408.

XX PR 03-AUG-2000; 2000US-00632366.

XX PR 21-SEP-2000; 2000US-0234687P.

XX PR 27-SEP-2000; 2000US-0236359P.

XX PR 04-OCT-2000; 2000GB-00024263.

XX PA (MOLE-) MOLECULAR DYNAMICS INC.

XX PI Penn SG, Hanzel DK, Chen W, Rank DR;

XX DR WPI; 2001-48897/53.

XX PT Human genome-derived single exon nucleic acid probes useful for analyzing  
 XX gene expression in human placenta.

XX PS Claim 27; SEQ ID NO 34186; 654pp; English.

XX The present invention relates to single exon nucleic acid probes (SENP;  
 CC see AAI31315-AAI57546). The present sequence is a peptide encoded by one  
 CC such probe. The probes are useful for producing a microarray for  
 CC predicting, measuring and displaying gene expression in samples derived  
 CC from human placenta. The probes are useful for antenatal diagnosis of  
 CC human genetic disorders

XX Sequence 30 AA;

Query Match 29.4%; Score 28.5; DB 4; Length 30;  
 Best Local Similarity 43.8%; Pred. No. 1.5e+03;  
 Matches 7; Conservative 4; Mismatches 4; Indels 1; Gaps 1;

Qy 3 NHLNSKIAFKIVSQEP 18  
 :|:|:  
 Db 13 NHRNLP-SFOIVTLDP 27

RESULT 198  
 AAM73727

ID AAM73727 standard; protein; 30 AA.

XX AC AAM73727;

DT 06-NOV-2001 (first entry)

XX Human bone marrow expressed probe encoded protein SEQ ID NO: 34033.

XX Human; bone marrow expressed exon; gene expression analysis; probe;  
 KW microarray; cancer; leukaemia; lymphoma; myeloma.

XX OS Homo sapiens.

XX PN WO200157276-A2.

XX PD 09-AUG-2001.

XX PF 30-JAN-2001; 2001WO-US000668.

XX PR 04-FEB-2000; 2000US-0180312P.

XX PR 26-MAY-2000; 2000US-0207456P.

XX PR 30-JUN-2000; 2000US-00608408.

XX PR 03-AUG-2000; 2000US-00632366.

XX PR 21-SEP-2000; 2000US-0234687P.

XX PR 27-SEP-2000; 2000US-0236359P.

XX PR 04-OCT-2000; 2000GB-00024263.

XX PA (MOLE-) MOLECULAR DYNAMICS INC.

XX PI Penn SG, Hanzel DK, Chen W, Rank DR;

XX DR WPI; 2001-488900/53.

XX PT Human genome-derived single exon nucleic acid probes useful for analyzing  
 XX gene expression in human bone marrow.

XX PS Example 4; SEQ ID NO 34033; 658pp + Sequence Listing; English.

XX The present invention provides a number of single exon nucleic acid  
 CC probes which are derived from genomic sequences expressed in the human  
 CC bone marrow. They can be used to measure gene expression in the human  
 CC samples, which may enable the improved diagnosis and treatment of cancers  
 CC such as lymphoma, leukaemia and myeloma. The present sequence is a  
 CC protein encoded by one of the probes of the invention

XX SQ Sequence 30 AA;

Query Match 29.4%; Score 28.5; DB 4; Length 30;  
 Best Local Similarity 43.8%; Pred. No. 1.5e+03;  
 Matches 7; Conservative 4; Mismatches 4; Indels 1; Gaps 1;

Qy 3 NHLNSKIAFKIVSQEP 18  
 :|:|:  
 Db 13 NHRNLP-SFOIVTLDP 27

RESULT 199  
 ABG55472

ID ABG55472 standard; peptide; 30 AA.

XX AC ABG55472;

DT 25-FEB-2003 (first entry)

XX Human liver peptide, SEQ ID No 34120.

XX Human; liver; cirrhosis; hyperlipoproteinaemia; hyperlipidaemia;  
 KW hypercholesterolaemia; coronary heart disease.

XX OS Homo sapiens.

```
XX PN WO200157273-A2.
XX PD
XX PF 09-AUG-2001.
XX PF 30-JAN-2001; 2001WO-US000664.
XX PR 04-FEB-2000; 2000US-0180312P.
XX PR 26-MAY-2000; 2000US-0207456P.
XX PR 30-JUN-2000; 2000US-00608408.
XX PR 03-AUG-2000; 2000US-00632366.
XX PR 21-SEP-2000; 2000US-0234687P.
XX PR 27-SEP-2000; 2000US-0236359P.
XX PR 04-OCT-2000; 2000GB-00024263.
XX PA (MOLE-) MOLECULAR DYNAMICS INC.
XX PI Penn SG, Hanzel DK, Chen W, Rank DR;
XX WPI; 2001-48898/53.
XX Human genome-derived single exon nucleic acid probes useful for analyzing
PT gene expression in human adult liver.
XX Claim 27; SEQ ID NO 34120; 658pp; English.
XX The invention relates to a single exon nucleic acid probe (SENP) (I) for
CC measuring human gene expression in a sample derived from human adult
CC liver, comprising one of 13109 defined nucleotide sequences given in the
CC specification (or complements/ fragments). The probe hybridises at high
CC stringency to a nucleic acid molecule expressed in the human adult liver.
CC (I) may be used for predicting, measuring and displaying gene expression
CC in samples derived from human adult liver. The genes identified may be
CC involved in genetic liver diseases such as cirrhosis,
CC hyperlipoproteinaemia, hyperlipidaemia and hypercholesterolaemia which is
CC associated with coronary heart disease. ABG47348-ABG59930 represent human
CC liver single exon encoded peptides of the invention. Note: The sequence
CC information for this patent does not appear in the printed specification
CC but was obtained in electronic format directly from WIPO at
CC ftp.wipo.int/pub/published_pct_sequences
XX Sequence 30 AA;
XX Query Match 29.4%; Score 28.5; DB 4; Length 30;
XX Best Local Similarity 43.8%; Pred. No. 1.5e+03;
XX Matches 7; Conservative 4; Mismatches 4; Indels 1; Gaps 1;
QY 3 NHLNKP-SQIVTLDP 18
DB 13 NHLNKP-SQIVTLDP 27
RESULT 200
ABG43609
ID ABG43609 standard; peptide; 30 AA.
AC ABG43609;
XX 19-AUG-2002 (first entry)
XX Human peptide encoded by genome-derived single exon probe SEQ ID 33274.
XX Human; single exon probe; asthma; lung cancer; COPD; IID;
XX Chronic obstructive pulmonary disease; interstitial lung disease;
XX familial idiopathic pulmonary fibrosis; neurofibromatosis;
XX tuberous sclerosis; Gaucher's disease; Niemann-Pick disease;
XX Hermansky-Pudlak syndrome; sarcoidosis; pulmonary haemosiderosis;
XX pulmonary histiocytosis; lymphangioleiomyomatosis; Karagenen syndrome;
XX pulmonary alveolar proteinosis; fibrocystic pulmonary dysplasia;
XX primary ciliary dyskinesia; pulmonary hypertension;
XX hyaline membrane disease.
XX Homo sapiens.
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XX PN WO200186003-A2.
XX PD
XX PF 15-NOV-2001.
XX PF 30-JAN-2001; 2001WO-US000665.
XX PR 04-FEB-2000; 2000US-0180312P.
XX PR 26-MAY-2000; 2000US-0207456P.
XX PR 30-JUN-2000; 2000US-00608408.
XX PR 03-AUG-2000; 2000US-00632366.
XX PR 21-SEP-2000; 2000US-0234687P.
XX PR 27-SEP-2000; 2000US-0236359P.
XX PR 04-OCT-2000; 2000GB-00024263.
XX PA (MOLE-) MOLECULAR DYNAMICS INC.
XX PI Penn SG, Hanzel DK, Chen W, Rank DR;
XX WPI; 2002-114183/15.
XX Spatially-addressable set of single exon nucleic acid probes, used to
PT measure gene expression in human lung samples.
XX Claim 27; SEQ ID NO 33274; 634pp; English.
XX The invention relates to a spatially-addressable set of single exon
CC nucleic acid probes for measuring gene expression in a sample derived
CC from human lung comprising single exon nucleic acid probes having one of
CC 12614 nucleic acid sequences mentioned in the specification, or their
CC complements or the 12387 open reading frames derived from the 12614
CC probes. Also included are a microarray comprising the novel set of probes
CC ; the novel set of probes which hybridise at high stringency to a nucleic
CC acid expressed in the human lung; measuring gene expression in a sample
CC derived from human lung, comprising (a) contacting the array with a
CC collection of detectably labeled nucleic acids derived from human lung
CC mRNA, and (b) measuring the label detectably bound to each probe of the
CC array; identifying exons in a eukaryotic genome, comprising (a)
CC algorithmically predicting at least one exon from genomic sequences of
CC the eukaryote; and (b) detecting specific hybridisation of detectably
CC labeled nucleic acids from eukaryote lung mRNA, to a single exon probe,
CC having a fragment identical to the predicted exon, the probe is included
CC in the above mentioned microarray; assigning exons to a single gene,
CC comprising (a) identifying exons from genomic sequence by the method
CC above and (b) measuring the expression of each of the exons in several
CC tissues and/or cell types using hybridisation to a single exon
CC microarrays having a probe with the exon, where a common pattern of
CC expression of the exons in the tissues and/or cell types indicates that
CC the exons should be assigned to a single gene; a peptide comprising one
CC of 12011 sequences, mentioned in the specification, or encoded by the
CC probes/open reading frames (ORF). The probes are used for gene expression
CC analysis, and for identifying exons in a gene, particularly using human
CC lung derived mRNA and for the study of lung diseases such as asthma, lung
CC cancer, chronic obstructive pulmonary disease (COPD), interstitial lung
CC disease (ILD), familial idiopathic pulmonary fibrosis, neurofibromatosis,
CC tuberous sclerosis, Gaucher's disease, Niemann-Pick disease, Hermansky-
CC Pudlak syndrome, sarcoidosis, pulmonary haemosiderosis, pulmonary
CC histiocytosis, lymphangioleiomyomatosis, pulmonary alveolar proteinosis,
CC Karagenen syndrome, fibrocystic pulmonary dysplasia, primary ciliary
CC dyskinesia, pulmonary hypertension and hyaline membrane disease. The
CC present sequence is a peptide/protein encoded by a single exon probe of
CC the invention. Note: The sequence data for this patent did not form part
CC of the printed specification, but was obtained in electronic format
CC directly from WIPO at ftp.wipo.int/pub/published_pct_sequences
XX Sequence 30 AA;
XX Query Match 29.4%; Score 28.5; DB 5; Length 30;
XX Best Local Similarity 43.8%; Pred. No. 1.5e+03;
XX Matches 7; Conservative 4; Mismatches 4; Indels 1; Gaps 1;
QY 3 NHLNKP-SQIVTLDP 18
XX 13 NHLNKP-SQIVTLDP 27
```



DD 13 NHKNLP-SFQIVTLDP 27

RESULT 201  
AAW27823

ID AAW27823 standard; protein; 42 AA.  
XX  
AC AAW27823;  
XX  
DT 21-JUL-1998 (first entry)  
XX  
DE Amino acid sequence of an ATP-dependent Clp proteinase chain clp.  
XX  
KW Staphylococcus aureus protein; ribozyme; antisense sequence; control;  
XX Staphylococcal gene; regulatory element; bacterial gene expression;  
KW vaccine; staphylococcal infection; food poisoning; scaled skin syndrome;  
KW toxic shock syndrome.  
XX  
OS Staphylococcus aureus.

XX  
XX Key Location/Qualifiers  
FH Misc-difference 1..42  
FT /note= "residues designated X are not defined in the  
FT specification"  
XX  
XX W09730070-A1.  
PN  
XX  
XX 21-AUG-1997. 97WO-US002318.  
PD  
XX  
PF 19-FEB-1997; 97WO-US002318.  
XX  
XX 20-FEB-1996; 96US-0011888P.  
PR  
XX (SMIK ) SMITHKLINE BEECHAM CORP.  
PA  
PI Black MT, Burnham MK, Hodgson JE, Knowles DJC, Nicholas RO;  
PI Pratt JM, Reichard RW, Rosenberg M, Ward JM;  
XX  
DR WI; 1997-424969/39.  
XX N-PSDB; AAT83788.  
XX  
XX Novel polypeptide(s) from Staphylococcus aureus strain WCUH29 - used to  
PT isolate antimicrobial compounds, and in vaccines against S. aureus  
PT infection.  
XX  
XX Claim 6; Page 294; 989pp; English.  
XX  
CC The present sequence represents a Staphylococcus aureus protein, that,  
CC based on homology with a Lactococcus protein, is believed to be an ATP-  
CC dependent Clp proteinase chain clp. The DNA sequence was isolated from a  
CC library of clones of S. aureus WCUH 29 in Escherichia coli. The DNA  
CC sequences can be used in the construction of ribozymes and antisense  
CC sequences to control the expression of Staphylococcal genes. The DNA  
CC sequence is also useful as a source of regulatory elements for the  
CC control of bacterial gene expression. The present protein may be used to  
CC produce vaccines to enable a host to produce specific antibodies with  
CC antibacterial action. These vaccines and antibodies would protect a host  
CC against invasion by S. aureus, and conditions relating to Staphylococcal  
CC infection, e.g. Staphylococcal food poisoning, scaled skin syndrome, and  
CC toxic shock syndrome  
XX  
SQ Sequence 42 AA;

Query Match 29.4%; Score 28.5; DB 2; Length 42;  
Best Local Similarity 50.0%; Pred. No. 2.2e+03;  
Matches 8; Conservative 1; Mismatches 4; Indels 3; Gaps 1;

QY 3 NHLNSKIAPKIVSQEP 18  
| | | | | :  
14 NLRLRSXI---IVKDQP 26  
| | | | | :

Db

RESULT 202

AAW78190  
ID AAW78190 standard; protein; 19 AA.  
XX  
AC AAW78190;  
XX  
DT 13-APR-1999 (first entry)  
XX  
DE Human secreted protein encoded by gene 65 clone HTDAG66.  
XX  
KW Human; secreted protein; fusion protein; gene therapy; protein therapy;  
KW diagnosis; tissue; cancer; tumour; neurodegenerative disorder; leukaemia;  
KW developmental abnormality; foetal deficiency; blood; allergy; renal;  
KW immune system; asthma; lymphocytic disease; brain; hepatic; lymphoma;  
KW inflammation; ischaemic shock; Alzheimer's disease; restenosis; AIDS;  
KW cognitive disorder; schizophrenia; prostate; obesity; osteoclast; thymus;  
KW osteoporosis; arthritis; testis; lung; thyroiditis; thyroid; digestion;  
KW endocrine; metabolism; regulation; malabsorption; gastritis; neoplasm.  
XX  
OS Homo sapiens.

XX  
XX Key Location/Qualifiers  
FH Misc-difference 19  
FT /label= unknown  
FT  
XX  
XX W09856804-A1.  
PN  
XX 17-DEC-1998.  
PD  
XX  
PF 11-JUN-1998; 98WO-US012125.  
XX  
XX 13-JUN-1997; 97US-0049547P.  
PR 13-JUN-1997; 97US-0049548P.  
PR 13-JUN-1997; 97US-0049549P.  
PR 13-JUN-1997; 97US-0049550P.  
PR 13-JUN-1997; 97US-0049566P.  
PR 13-JUN-1997; 97US-0049606P.  
PR 13-JUN-1997; 97US-0049607P.  
PR 13-JUN-1997; 97US-0049608P.  
PR 13-JUN-1997; 97US-0049609P.  
PR 13-JUN-1997; 97US-0049610P.  
PR 13-JUN-1997; 97US-0049611P.  
PR 13-JUN-1997; 97US-0050901P.  
PR 13-JUN-1997; 97US-0052989P.  
PR 08-JUL-1997; 97US-0051919P.  
PR 18-AUG-1997; 97US-0055984P.  
PR 12-SEP-1997; 97US-0058665P.  
PR 12-SEP-1997; 97US-0058668P.  
PR 12-SEP-1997; 97US-0058669P.  
PR 12-SEP-1997; 97US-0058750P.  
PR 12-SEP-1997; 97US-0058971P.  
PR 12-SEP-1997; 97US-0058972P.  
PR 12-SEP-1997; 97US-0058975P.  
PR 02-OCT-1997; 97US-0060834P.  
PR 02-OCT-1997; 97US-0060841P.  
PR 02-OCT-1997; 97US-0060844P.  
PR 02-OCT-1997; 97US-0060865P.  
PR 02-OCT-1997; 97US-0061059P.  
PR 02-OCT-1997; 97US-0061060P.  
XX  
PA (HUMA-) HUMAN GENOME SCI INC.  
XX  
XX Moore PA, Shi Y, Rosen CA, Ruben SM, Lafleur DW, Olsen HS;  
PI Ebner R, Brewer LA, Young P, Greene JM, Ferrie AM, Yu G, Ni J;  
PI Peng P;  
XX  
XX WPI; 1999-080881/07.  
DR N-PSDB; AAX04375.  
XX  
XX New isolated human genes and the secreted polypeptides they encode -  
PT useful for diagnosis and treatment of e.g. cancers, neurological  
PT disorders, immune diseases, inflammation or blood disorders.  
XX  
XX Claim 11; Page 300-301; 380pp; English.  
PS

XX CC This sequence represents a secreted human protein encoded by the gene  
 CC clone detailed in the descriptor line. The gene can be used to generate  
 CC fusion proteins by linking to the gene to a human immunoglobulin Fc  
 CC portion (e.g. AAX04302) for increasing the stability of the fused protein  
 CC as compared to the human protein only. The invention relates to 86 novel  
 CC genes and their fragments (nucleic acid sequences: AAX04311-X04410; amino  
 CC acid sequences AAW78126-W78225) which are useful for preventing, treating  
 CC or ameliorating medical conditions e.g. by protein or gene therapy. Also,  
 CC pathological conditions can be diagnosed by determining the amount of the  
 CC new polypeptides in a sample or by determining the presence of mutations  
 CC in the new polynucleotides. Specific uses are described for each of the  
 CC 86 polynucleotides, based on which tissues they are most highly expressed  
 CC in (see AAX04311 for described uses)

XX SQ Sequence 19 AA;  
 Query Match 28.9%; Score 28; DB 2; Length 19;  
 Best Local Similarity 62.5%; Pred. No. 1e+03;  
 Matches 5; Conservative 0; Mismatches 3; Indels 0; Gaps 0;

OY 2 PNHLNSKI 9  
 DB 5 PTHLEGI 12

RESULT 203  
 ABB40664  
 ID ABB40664 standard; peptide; 19 AA.  
 XX AC ABB40664;  
 XX DT 04-FEB-2002 (first entry)  
 DE Peptide #8170 encoded by human foetal liver single exon probe.  
 XX Human; foetal liver; gene expression; single exon nucleic acid probe.  
 XX Homo sapiens.  
 XX WO200157277-A2.  
 XX 09-AUG-2001.  
 XX 30-JAN-2001; 2001WO-US000669.  
 XX 04-FEB-2000; 2000US-0180312P.  
 XX 26-MAY-2000; 2000US-0207456P.  
 XX 30-JUN-2000; 2000US-00608408.  
 XX 03-AUG-2000; 2000US-00632366.  
 XX 21-SEP-2000; 2000US-0234687P.  
 XX 27-SEP-2000; 2000US-0236359P.  
 XX 04-OCT-2000; 2000GB-00024263.  
 XX (MOLE-) MOLECULAR DYNAMICS INC.  
 XX Penn SG, Hanzel DK, Chen W, Rank DR;  
 XX WPI; 2001-483447/52.  
 XX Human genome-derived single exon nucleic acid probes useful for analyzing  
 XX gene expression in human fetal liver.  
 XX Claim 27; SEQ ID NO 33299; 639pp + Sequence Listing; English.  
 XX The invention relates to a single exon nucleic acid probe for measuring  
 XX human gene expression in a sample derived from human foetal liver. The  
 XX single exon nucleic acid probes may be used for predicting, measuring and  
 XX displaying gene expression in samples derived from human fetal liver. The  
 XX present sequence is a peptide encoded by a single exon nucleic acid probe  
 XX of the invention. Note: The sequence data for this patent did not form  
 XX part of the printed specification, but was obtained in electronic format  
 XX directly from WIPO at ftp.wipo.int/pub/published\_pct\_sequences

XX SQ Sequence 19 AA;  
 Query Match 28.9%; Score 28; DB 4; Length 19;  
 Best Local Similarity 50.0%; Pred. No. 1e+03;  
 Matches 6; Conservative 2; Mismatches 4; Indels 0; Gaps 0;

OY 6 NSKIAFKIVSOE 17  
 DB 6 NYKLSVKFSSOE 17

RESULT 204  
 AAM34425  
 ID AAM34425 standard; protein; 19 AA.  
 XX AC AAM34425;  
 XX DT 17-OCT-2001 (first entry)  
 DE Peptide #8462 encoded by probe for measuring placental gene expression.  
 XX Probe; microarray; human; placenta; antenatal diagnosis;  
 XX genetic disorder.  
 XX Homo sapiens.  
 XX WO200157272-A2.  
 XX 09-AUG-2001.  
 XX 30-JAN-2001; 2001WO-US000663.  
 XX 04-FEB-2000; 2000US-0180312P.  
 XX 26-MAY-2000; 2000US-0207456P.  
 XX 30-JUN-2000; 2000US-00608408.  
 XX 03-AUG-2000; 2000US-00632366.  
 XX 21-SEP-2000; 2000US-0234687P.  
 XX 27-SEP-2000; 2000US-0236359P.  
 XX 04-OCT-2000; 2000GB-00024263.  
 XX (MOLE-) MOLECULAR DYNAMICS INC.  
 XX Penn SG, Hanzel DK, Chen W, Rank DR;  
 XX WPI; 2001-488897/53.  
 XX Human genome-derived single exon nucleic acid probes useful for analyzing  
 XX gene expression in human placenta.  
 XX Claim 27; SEQ ID NO 34694; 654pp; English.  
 XX The present invention relates to single exon nucleic acid probes (SENP;  
 XX see AAI31315-AAI57546). The present sequence is a peptide encoded by one  
 XX such probe. The probes are useful for producing a microarray for  
 XX predicting, measuring and displaying gene expression in samples derived  
 XX from human placenta. The probes are useful for antenatal diagnosis of  
 XX human genetic disorders

XX SQ Sequence 19 AA;  
 Query Match 28.9%; Score 28; DB 4; Length 19;  
 Best Local Similarity 50.0%; Pred. No. 1e+03;  
 Matches 6; Conservative 2; Mismatches 4; Indels 0; Gaps 0;

OY 6 NSKIAFKIVSOE 17  
 DB 6 NYKLSVKFSSOE 17

RESULT 205  
 AAM74313  
 ID AAM74313 standard; protein; 19 AA.

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XX AC AAM74313;
XX DT 06-NOV-2001 (first entry)
XX DE Human bone marrow expressed probe encoded protein SEQ ID NO: 34619.
XX KW Human; bone marrow expressed exon; gene expression analysis; probe;
XX KW microarray; cancer; leukaemia; lymphoma; myeloma.
XX OS Homo sapiens.
XX PN WO200157276-A2.
XX PD 09-AUG-2001.
XX PF 30-JAN-2001; 2001WO-US000668.
XX PR 04-FEB-2000; 2000US-0180312P.
XX PR 26-MAY-2000; 2000US-0207456P.
XX PR 30-JUN-2000; 2000US-00608408.
XX PR 03-AUG-2000; 2000US-00632366.
XX PR 21-SEP-2000; 2000US-0234687P.
XX PR 27-SEP-2000; 2000US-0236359P.
XX PR 04-OCT-2000; 2000GB-00024263.
XX PA (MOLE-) MOLECULAR DYNAMICS INC.
XX PI Penn SG, Hanzel DK, Chen W, Rank DR;
XX DR WPI; 2001-488900/53.
XX PT Human genome-derived single exon nucleic acid probes useful for analyzing
XX PT gene expression in human bone marrow.
XX PS Example 4; SEQ ID NO 34619; 658pp + Sequence Listing; English.
XX CC The present invention provides a number of single exon nucleic acid
XX CC probes which are derived from genomic sequences expressed in the human
XX CC bone marrow. They can be used to measure gene expression in bone marrow
XX CC samples, which may enable the improved diagnosis and treatment of cancers
XX CC such as lymphoma, leukaemia and myeloma. The present sequence is a
XX CC protein encoded by one of the probes of the invention
XX SQ Sequence 19 AA;
XX Query Match 28.9%; Score 28; DB 4; Length 19;
XX Best Local Similarity 50.0%; Pred. No. 1e+03;
XX Matches 6; Conservative 2; Mismatches 4; Indels 0; Gaps 0;
XX Qy 6 NSKIAFKIVSQE 17
XX Db 6 NYKLSVRFSSQE 17
XX RESULT 206
XX ID AAM61524 standard; protein; 19 AA.
XX AC AAM61524;
XX DT 05-NOV-2001 (first entry)
XX DE Human brain expressed single exon probe encoded protein SEQ ID NO: 33629.
XX KW Human; brain expressed exon; gene expression analysis; probe; microarray;
XX KW Alzheimer's disease; multiple sclerosis; schizophrenia; epilepsy; cancer.
XX OS Homo sapiens.
XX PN WO200157275-A2.
XX PD 09-AUG-2001.
XX Query Match 28.9%; Score 28; DB 4; Length 19;
XX Best Local Similarity 50.0%; Pred. No. 1e+03;
XX Matches 6; Conservative 2; Mismatches 4; Indels 0; Gaps 0;
XX Qy 6 NSKIAFKIVSQE 17
XX Db 6 NYKLSVRFSSQE 17
XX RESULT 207
XX ID ABG56112 standard; peptide; 19 AA.
XX AC ABG56112;
XX DT 25-FEB-2003 (first entry)
XX DE Human liver peptide, SEQ ID No 34760.
XX KW Human; liver; cirrhosis; hyperlipoproteinaemia; hyperlipidaemia;
XX KW hypercholesterolaemia; coronary heart disease.
XX OS Homo sapiens.
XX PN WO200157273-A2.
XX PD 09-AUG-2001.
XX PF 30-JAN-2001; 2001WO-US000664.
XX PR 04-FEB-2000; 2000US-0180312P.
XX PR 26-MAY-2000; 2000US-0207456P.
XX PR 30-JUN-2000; 2000US-00608408.
XX PR 03-AUG-2000; 2000US-00632366.
XX PR 21-SEP-2000; 2000US-0234687P.
XX PR 27-SEP-2000; 2000US-0236359P.
XX PR 04-OCT-2000; 2000GB-00024263.
XX PA (MOLE-) MOLECULAR DYNAMICS INC.
XX PI Penn SG, Hanzel DK, Chen W, Rank DR;

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XX PF 30-JAN-2001; 2001WO-US000667.
XX PR 04-FEB-2000; 2000US-0180312P.
XX PR 26-MAY-2000; 2000US-0207456P.
XX PR 30-JUN-2000; 2000US-00608408.
XX PR 03-AUG-2000; 2000US-00632366.
XX PR 21-SEP-2000; 2000US-0234687P.
XX PR 27-SEP-2000; 2000US-0236359P.
XX PR 04-OCT-2000; 2000GB-00024263.
XX PA (MOLE-) MOLECULAR DYNAMICS INC.
XX PI Penn SG, Hanzel DK, Chen W, Rank DR;
XX DR WPI; 2001-483446/52.
XX PT Single exon nucleic acid probes for analyzing gene expression in human
XX PT brains.
XX PS Example 4; SEQ ID NO 33629; 650pp + Sequence Listing; English.
XX CC The present invention provides a number of single exon nucleic acid
XX CC probes which are derived from genomic sequences expressed in the human
XX CC brain. They can be used to measure gene expression in brain cell samples,
XX CC which may enable the diagnosis and improved treatment of nervous system
XX CC diseases such as Alzheimer's disease, multiple sclerosis, schizophrenia,
XX CC epilepsy and cancers. The present sequence is a protein encoded by one of
XX CC the probes of the invention
XX SQ Sequence 19 AA;
XX Query Match 28.9%; Score 28; DB 4; Length 19;
XX Best Local Similarity 50.0%; Pred. No. 1e+03;
XX Matches 6; Conservative 2; Mismatches 4; Indels 0; Gaps 0;
XX Qy 6 NSKIAFKIVSQE 17
XX Db 6 NYKLSVRFSSQE 17
XX RESULT 207
XX ID ABG56112 standard; peptide; 19 AA.
XX AC ABG56112;
XX DT 25-FEB-2003 (first entry)
XX DE Human liver peptide, SEQ ID No 34760.
XX KW Human; liver; cirrhosis; hyperlipoproteinaemia; hyperlipidaemia;
XX KW hypercholesterolaemia; coronary heart disease.
XX OS Homo sapiens.
XX PN WO200157273-A2.
XX PD 09-AUG-2001.
XX PF 30-JAN-2001; 2001WO-US000664.
XX PR 04-FEB-2000; 2000US-0180312P.
XX PR 26-MAY-2000; 2000US-0207456P.
XX PR 30-JUN-2000; 2000US-00608408.
XX PR 03-AUG-2000; 2000US-00632366.
XX PR 21-SEP-2000; 2000US-0234687P.
XX PR 27-SEP-2000; 2000US-0236359P.
XX PR 04-OCT-2000; 2000GB-00024263.
XX PA (MOLE-) MOLECULAR DYNAMICS INC.
XX PI Penn SG, Hanzel DK, Chen W, Rank DR;

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PN WO9931117-A1.
XX
PD 24-JUN-1999.
XX
PF 17-DEC-1998; 98WO-US027059.
XX
PR 18-DEC-1997; 97US-0068006P.
PR 18-DEC-1997; 97US-0068007P.
PR 18-DEC-1997; 97US-0068008P.
PR 18-DEC-1997; 97US-0068053P.
PR 18-DEC-1997; 97US-0068057P.
PR 18-DEC-1997; 97US-0068054P.
PR 18-DEC-1997; 97US-0068064P.
PR 18-DEC-1997; 97US-0070923P.
PR 19-DEC-1997; 97US-0068163P.
PR 19-DEC-1997; 97US-0068365P.
PR 19-DEC-1997; 97US-0068367P.
PR 19-DEC-1997; 97US-0068368P.
PR 19-DEC-1997; 97US-0068369P.
XX
PA (HUMA-) HUMAN GENOME SCI INC.
XX
PI Moore PA, Ruben SM, Carter KC, Shi Y, Rosen CA, Soppet DR;
PI Kyaw H, Wei Y, Florence K, Duan RD, Florence C, Greene JM, Feng P;
PI Ferrie AM, Yu G, Janat F, Ni J;
XX
DR WPI; 1999-418749/35.
XX
XX New isolated human genes encoding secreted polypeptides.
PT Disclosure; Page 312; 537pp; English.
PS
XX
CC AAX97916 to AAX98029 represent 110 isolated human secreted protein genes.
CC AAX36224 to AAX36727 represent the secreted proteins encoded by the 110
CC human genes. The genes and their corresponding secreted polypeptides are
CC useful for preventing, treating or ameliorating medical conditions, e.g.
CC by protein or gene therapy. Also pathological conditions can be diagnosed
CC by determining the amount of the new polypeptides in a sample or by
CC determining the presence of mutations in the new genes. Specific uses are
CC described for each of the 110 genes, based on which tissues they are most
CC highly expressed in, and include developing products for the diagnosis or
CC treatment of cancer, tumours, developmental abnormalities and foetal
CC deficiencies, blood disorders, diseases of the immune system, autoimmune
CC diseases, inflammation, allergies, Alzheimer's and cognitive disorders,
CC schizophrenia, arthritis, asthma, psoriasis, sepsis, skin disorders,
CC atherosclerosis, diabetes, cardiovascular disorders, kidney disorders,
CC digestive/endocrine disorders, infections and AIDS. The polypeptides are
CC also useful for identifying their binding partners. The sequences given
CC in AAX97907 to AAX97915 and AAX36223 are used in the exemplification of
CC the present invention
XX
XX Sequence 23 AA;
SQ
Query Match 28.9%; Score 28; DB 2; Length 23;
Best Local Similarity 27.8%; Pred. No. 1.3e+03;
Matches 5; Conservative 7; Mismatches 6; Indels 0; Gaps 0;
QY 1 EPNHLSKIAPKIVSQEP 18
DB 5 EPQCGASRLSWKLNSSP 22
||| :|:|:|:|:|
||| :|:|:|:|:|

RESULT 212
AAM15118
ID AAM15118 standard; protein; 23 AA.
XX
AC AAM15118;
XX
DT 12-OCT-2001 (first entry)
XX
DE Peptide #1552 encoded by probe for measuring cervical gene expression.
XX
PR Probe; human; microarray; gene expression; cervical epithelial cell;
KW

KW cervical cancer.
XX
OS Homo sapiens.
XX
PN WO200157278-A2.
XX
PD 09-AUG-2001.
XX
PF 30-JAN-2001; 2001WO-US000670.
XX
PR 04-FEB-2000; 2000US-0180312P.
PR 26-MAY-2000; 2000US-0207456P.
PR 30-JUN-2000; 2000US-00608408.
PR 03-AUG-2000; 2000US-00632366.
PR 21-SEP-2000; 2000US-0234687P.
PR 27-SEP-2000; 2000US-0236359P.
PR 04-OCT-2000; 2000GB-00024263.
XX
PA (MOLE-) MOLECULAR DYNAMICS INC.
XX
PI Penn SG, Hanzel DK, Chen W, Rank DR;
XX
XX WPI; 2001-489901/53.
XX
XX Human genome-derived single exon nucleic acid probes useful for analyzing
XX gene expression in human cervical epithelial cells.
XX
XX Claim 27; SEQ ID NO 19944; 487pp; English.
XX
XX The present invention relates to human single exon nucleic acid probes
XX (SENPs: see AAI10068-AAI28459). The present sequence is a peptide encoded
XX by one such probe. The SENPs are derived from human HeLa cells. The SENPs
XX can be used to produce a single exon microarray, which can be used for
XX measuring human gene expression in a sample derived from human cervical
XX epithelial cells. By measuring gene expression, the probes are therefore
XX useful in grading and/or staging of diseases of the cervix, notably
XX cervical cancer. Note: The sequence data for this patent did not form
XX part of the printed specification, but was obtained in electronic format
XX directly from WIPO at ftp.wipo.int/pub/published_pct_sequences
XX
XX Sequence 23 AA;
SQ
Query Match 28.9%; Score 28; DB 4; Length 23;
Best Local Similarity 42.9%; Pred. No. 1.3e+03;
Matches 6; Conservative 3; Mismatches 5; Indels 0; Gaps 0;
QY 5 LNSKIAPKIVSQEP 18
DB 4 LKSFSLSIQISKQEP 17
||| :|:|:|:|:|
||| :|:|:|:|:|

RESULT 213
ABB34112
ID ABB34112 standard; peptide; 23 AA.
XX
AC ABB34112;
XX
DT 04-FEB-2002 (first entry)
XX
XX Peptide #1618 encoded by human foetal liver single exon probe.
XX
KW Human; foetal liver; gene expression; single exon nucleic acid probe.
XX
OS Homo sapiens.
XX
PN WO200157277-A2.
XX
PD 09-AUG-2001.
XX
PF 30-JAN-2001; 2001WO-US000669.
XX
PR 04-FEB-2000; 2000US-0180312P.
PR 26-MAY-2000; 2000US-0207456P.
XX
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PR 30-JUN-2000; 2000US-00608408.  
 PR 03-AUG-2000; 2000US-00632366.  
 PR 21-SEP-2000; 2000US-0234687P.  
 PR 27-SEP-2000; 2000US-0236359P.  
 PR 04-OCT-2000; 2000GB-00024263.  
 XX (MOLE-) MOLECULAR DYNAMICS INC.  
 PA Penn SG, Hanzel DK, Chen W, Rank DR;  
 PI WPI; 2001-483447/52.  
 XX Human genome-derived single exon nucleic acid probes useful for analyzing  
 PT gene expression in human fetal liver.  
 XX  
 PS Claim 27; SEQ ID NO 26747; 639pp + Sequence Listing; English.  
 XX The invention relates to a single exon nucleic acid probe for measuring  
 CC human gene expression in a sample derived from human foetal liver. The  
 CC single exon nucleic acid probes may be used for predicting, measuring and  
 CC displaying gene expression in samples derived from human fetal liver. The  
 CC present sequence is a peptide encoded by a single exon nucleic acid probe  
 CC of the invention. Note: The sequence data for this patent did not form  
 CC part of the printed specification, but was obtained in electronic format  
 CC directly from WIPO at ftp.wipo.int/pub/published\_pct\_sequences  
 CC  
 XX Sequence 23 AA;  
 SQ  
 Query Match 28.9%; Score 28; DB 4; Length 23;  
 Best Local Similarity 42.9%; Pred. No. 1.3e+03;  
 Matches 6; Conservative 3; Mismatches 5; Indels 0; Gaps 0;  
 QY 5 LNSKIAFKIVSQEP 18  
 Db 4 LKSFSLSIQISKQEP 17  
 RESULT 214  
 AAM27573  
 ID AAM27573 standard; protein; 23 AA.  
 XX  
 AC AAM27573;  
 XX 17-OCT-2001 (first entry)  
 XX Peptide #1610 encoded by probe for measuring placental gene expression.  
 DE Probe; microarray; human; placenta; antenatal diagnosis;  
 XX genetic disorder.  
 KW Homo sapiens.  
 XX  
 OS WO200157272-A2.  
 XX  
 PN 09-AUG-2001.  
 XX  
 PD 30-JAN-2001; 2001WO-US000663.  
 XX  
 PF 04-FEB-2000; 2000US-0180312P.  
 XX  
 PR 26-MAY-2000; 2000US-0207456P.  
 PR 30-JUN-2000; 2000US-00608408.  
 PR 03-AUG-2000; 2000US-00632366.  
 PR 21-SEP-2000; 2000US-0234687P.  
 PR 27-SEP-2000; 2000US-0236359P.  
 PR 04-OCT-2000; 2000GB-00024263.  
 XX (MOLE-) MOLECULAR DYNAMICS INC.  
 PA Penn SG, Hanzel DK, Chen W, Rank DR;  
 PI WPI; 2001-488897/53.  
 XX Human genome-derived single exon nucleic acid probes useful for analyzing  
 PT

PT gene expression in human placenta.  
 XX Claim 27; SEQ ID NO 27842; 654pp; English.  
 XX The present invention relates to single exon nucleic acid probes (SENP:  
 CC see AAT31315-AA157546). The present sequence is a peptide encoded by one  
 CC such probe. The probes are useful for producing a microarray for  
 CC predicting, measuring and displaying gene expression in samples derived  
 CC from human placenta. The probes are useful for antenatal diagnosis of  
 CC human genetic disorders  
 XX Sequence 23 AA;  
 SQ  
 Query Match 28.9%; Score 28; DB 4; Length 23;  
 Best Local Similarity 42.9%; Pred. No. 1.3e+03;  
 Matches 6; Conservative 3; Mismatches 5; Indels 0; Gaps 0;  
 QY 5 LNSKIAFKIVSQEP 18  
 Db 4 LKSFSLSIQISKQEP 17  
 RESULT 215  
 ABB28942  
 ID ABB28942 standard; peptide; 23 AA.  
 XX  
 AC ABB28942;  
 XX 01-FEB-2002 (first entry)  
 XX Peptide #1593 encoded by breast cell single exon nucleic acid probe.  
 DE Human; microarray; single exon probe; gene expression; breast; disease;  
 XX cancer.  
 KW Homo sapiens.  
 XX  
 OS WO200157271-A2.  
 XX  
 PN 09-AUG-2001.  
 XX  
 PD 30-JAN-2001; 2001WO-US000662.  
 XX  
 PF 04-FEB-2000; 2000US-0180312P.  
 XX  
 PR 26-MAY-2000; 2000US-0207456P.  
 PR 30-JUN-2000; 2000US-00608408.  
 PR 03-AUG-2000; 2000US-00632366.  
 PR 21-SEP-2000; 2000US-0234687P.  
 PR 27-SEP-2000; 2000US-0236359P.  
 PR 04-OCT-2000; 2000GB-00024263.  
 XX (MOLE-) MOLECULAR DYNAMICS INC.  
 PA Penn SG, Hanzel DK, Chen W, Rank DR;  
 PI WPI; 2001-496933/54.  
 XX New spatially-addressable set of single exon nucleic acid probes, useful  
 PT for measuring gene expression in sample derived from human breast,  
 PT comprises number of single exon nucleic acid probes.  
 XX Claim 27; SEQ ID NO 11910; 327pp + Sequence Listing; English.  
 XX The invention relates to a spatially-addressable set of single exon  
 CC nucleic acid probes for measuring gene expression in a sample derived  
 CC from human breast and BT 474 cells. The method involves contacting the  
 CC probes with a collection of detectably labelled nucleic acids derived  
 CC from mRNA of human breast, and then measuring the label bound to each  
 CC probe of the microarray. The probes are useful for verifying the  
 CC expression of regions of genomic DNA predicted to encode proteins. They  
 CC are useful for gene discovery, and for determining predisposition and/or  
 CC prognosing breast disease. Gene expression analysis is useful for  
 CC assessing the toxicity of chemical agents on cells. The microarray of

CC this invention presents a far greater diversity of probes for measuring  
CC gene expression, with far less bias than expressed sequence tag  
CC microarrays. The method is suitable for rapid production of functional  
CC information from genomic sequence. The present sequence is a peptide  
CC encoded by a single exon nucleic acid probe of the invention. Note: The  
CC sequence data for this patent did not form part of the printed  
CC specification, but was obtained in electronic format directly from WIPO  
CC at ftp.wipo.int/pub/published\_pct\_sequences

XX SQ Sequence 23 AA;  
Query Match 28.9%; Score 28; DB 4; Length 23;  
Best Local Similarity 42.9%; Pred. No. 1.3e+03;  
Matches 6; Conservative 3; Mismatches 5; Indels 0; Gaps 0;

QY 5 LNSKIAFKIVSQEP 18  
Db 4 LKSFSLIQISKQEP 17

RESULT 216  
ABBI19553  
ID ABBI19553 standard; protein; 23 AA.  
XX  
AC ABBI19553;  
XX  
DT 23-JAN-2002 (first entry)  
XX  
DE Protein #1552 encoded by probe for measuring heart cell gene expression.  
XX  
KW Human; gene expression; heart; microarray; vascular system;  
KW cardiovascular disease; hypertension; cardiac arrhythmia;  
KW congenital heart disease.  
XX  
OS Homo sapiens.  
XX  
PN WO200157274-A2.  
XX  
PD 09-AUG-2001.  
XX  
PF 30-JAN-2001; 2001WO-US000666.  
XX  
PR 04-FEB-2000; 2000US-0180312P.  
PR 26-MAY-2000; 2000US-0207456P.  
PR 30-JUN-2000; 2000US-00608408.  
PR 03-AUG-2000; 2000US-00632366.  
PR 21-SEP-2000; 2000US-0234687P.  
PR 27-SEP-2000; 2000US-0236359P.  
PR 04-OCT-2000; 2000GB-00024263.  
XX  
PA (MOLE-) MOLECULAR DYNAMICS INC.  
XX  
PI Penn SG, Hanzel DK, Chen W, Rank DR;  
XX  
DR WPI; 2001-488990/53.  
XX  
PT Single exon nucleic acid probes for analyzing gene expression in human  
PT hearts.  
XX  
PS Claim 15; SEQ ID NO 21323; 530pp; English.  
XX  
CC The present invention relates to single exon nucleic acid probes for  
CC measuring human gene expression in a sample derived from human heart (see  
CC ABA21535-ABA41305). The present sequence is a protein encoded by one such  
CC probe. The probes may be used for predicting, measuring and displaying  
CC gene expression in samples derived from the human heart via microarrays.  
CC By measuring gene expression, the probes are useful for predicting,  
CC diagnosing, grading, staging, monitoring and prognosing diseases of the  
CC human heart and vascular system e.g. cardiovascular disease,  
CC hypertension, cardiac arrhythmias and congenital heart disease. Note: The  
CC sequence data for this patent did not form part of the printed  
CC specification, but was obtained in electronic format directly from WIPO  
CC at ftp.wipo.int/pub/published\_pct\_sequences

XX SQ Sequence 23 AA;  
Query Match 28.9%; Score 28; DB 4; Length 23;  
Best Local Similarity 42.9%; Pred. No. 1.3e+03;  
Matches 6; Conservative 3; Mismatches 5; Indels 0; Gaps 0;

QY 5 LNSKIAFKIVSQEP 18  
Db 4 LKSFSLIQISKQEP 17

RESULT 217  
AAM67281  
ID AAM67281 standard; protein; 23 AA.  
XX  
AC AAM67281;  
XX  
DT 06-NOV-2001 (first entry)  
XX  
DE Human bone marrow expressed probe encoded protein SEQ ID NO: 27587.  
XX  
KW Human; bone marrow expressed exon; gene expression analysis; probe;  
KW microarray; cancer; leukaemia; lymphoma; myeloma.  
XX  
OS Homo sapiens.  
XX  
PN WO200157276-A2.  
XX  
PD 09-AUG-2001.  
XX  
PF 30-JAN-2001; 2001WO-US000668.  
XX  
PR 04-FEB-2000; 2000US-0180312P.  
PR 26-MAY-2000; 2000US-0207456P.  
PR 30-JUN-2000; 2000US-00608408.  
PR 03-AUG-2000; 2000US-00632366.  
PR 21-SEP-2000; 2000US-0234687P.  
PR 27-SEP-2000; 2000US-0236359P.  
PR 04-OCT-2000; 2000GB-00024263.  
XX  
PA (MOLE-) MOLECULAR DYNAMICS INC.  
XX  
PI Penn SG, Hanzel DK, Chen W, Rank DR;  
XX  
DR WPI; 2001-488990/53.  
XX  
PT Human genome-derived single exon nucleic acid probes useful for analyzing  
PT gene expression in human bone marrow.  
XX  
PS Example 4; SEQ ID NO 27587; 658pp + Sequence Listing; English.  
XX  
CC The present invention provides a number of single exon nucleic acid  
CC probes which are derived from genomic sequences expressed in the human  
CC bone marrow. They can be used to measure gene expression in bone marrow  
CC samples, which may enable the improved diagnosis and treatment of cancers  
CC such as lymphoma, leukaemia and myeloma. The present sequence is a  
CC protein encoded by one of the probes of the invention  
XX  
XX SQ Sequence 23 AA;  
Query Match 28.9%; Score 28; DB 4; Length 23;  
Best Local Similarity 42.9%; Pred. No. 1.3e+03;  
Matches 6; Conservative 3; Mismatches 5; Indels 0; Gaps 0;

QY 5 LNSKIAFKIVSQEP 18  
Db 4 LKSFSLIQISKQEP 17

RESULT 218  
AAM54900  
ID AAM54900 standard; protein; 23 AA.



XX AAM54900;  
AC  
XX  
DT 05-NOV-2001 (first entry)  
XX  
DE Human brain expressed single exon probe encoded protein SEQ ID NO: 27005.  
XX  
KW Human; brain expressed exon; gene expression analysis; probe; microarray;  
KW Alzheimer's disease; multiple sclerosis; schizophrenia; epilepsy; cancer.  
XX  
OS Homo sapiens.  
XX  
PN WO200157275-A2.  
XX  
PD 09-AUG-2001.  
XX  
PF 30-JAN-2001; 2001WO-US000667.  
XX  
PR 04-FEB-2000; 2000US-0180312P.  
PR 26-MAY-2000; 2000US-0207456P.  
PR 30-JUN-2000; 2000US-00608408.  
PR 03-AUG-2000; 2000US-00632366.  
PR 21-SEP-2000; 2000US-0234687P.  
PR 27-SEP-2000; 2000US-0236359P.  
PR 04-OCT-2000; 2000GB-00024263.  
XX  
PA (MOLE-) MOLECULAR DYNAMICS INC.  
XX  
PI Penn SG, Hanzel DK, Chen W, Rank DR;  
XX  
DR WPI; 2001-483446/52.  
XX  
PT Single exon nucleic acid probes for analyzing gene expression in human  
PT brains.  
XX  
XX Example 4; SEQ ID NO 27005; 650pp + Sequence Listing; English.  
PS  
SS The present invention provides a number of single exon nucleic acid  
CC probes which are derived from genomic sequences expressed in the human  
CC brain. They can be used to measure gene expression in brain cell samples,  
CC which may enable the diagnosis and improved treatment of nervous system  
CC diseases such as Alzheimer's disease, multiple sclerosis, schizophrenia,  
CC epilepsy and cancers. The present sequence is a protein encoded by one of  
CC the probes of the invention  
XX  
SQ Sequence 23 AA;  
Query Match 28.9%; Score 28; DB 4; Length 23;  
Best Local Similarity 42.9%; Pred. No. 1.3e+03;  
Matches 6; Conservative 3; Mismatches 5; Indels 0; Gaps 0;  
QY 5 LNSKIAPKIVSQEP 18  
Db | : : : |||  
4 LKSFLSIQISKQEP 17  
RESULT 219  
ABG48943  
ID ABG48943 standard; peptide; 23 AA.  
XX  
AC ABG48943;  
XX  
DT 25-FEB-2003 (first entry)  
XX  
DE Human liver peptide, SEQ ID No 27591.  
XX  
KW Human; liver; cirrhosis; hyperlipoproteinaemia; hyperlipidaemia;  
KW hypercholesterolaemia; coronary heart disease.  
XX  
OS Homo sapiens.  
XX  
PN WO200157273-A2.  
XX

PD 09-AUG-2001.  
XX  
PF 30-JAN-2001; 2001WO-US000664.  
XX  
PR 04-FEB-2000; 2000US-0180312P.  
PR 26-MAY-2000; 2000US-0207456P.  
PR 30-JUN-2000; 2000US-00608408.  
PR 03-AUG-2000; 2000US-00632366.  
PR 21-SEP-2000; 2000US-0234687P.  
PR 27-SEP-2000; 2000US-0236359P.  
PR 04-OCT-2000; 2000GB-00024263.  
XX  
PA (MOLE-) MOLECULAR DYNAMICS INC.  
XX  
PI Penn SG, Hanzel DK, Chen W, Rank DR;  
XX  
DR WPI; 2001-488898/53.  
XX  
PT Human genome-derived single exon nucleic acid probes useful for analyzing  
PT gene expression in human adult liver.  
XX  
XX Claim 27; SEQ ID NO 27591; 658pp; English.  
XX  
CC The invention relates to a single exon nucleic acid probe (SENP) (I) for  
CC measuring human gene expression in a sample derived from human adult  
CC liver, comprising one of 13109 defined nucleotide sequences given in the  
CC specification (or complements/ fragments). The probe hybridises at high  
CC stringency to a nucleic acid molecule expressed in the human adult liver.  
CC (I) may be used for predicting, measuring and displaying gene expression  
CC in samples derived from human adult liver. The genes identified may be  
CC involved in genetic liver diseases such as cirrhosis,  
CC hyperlipoproteinaemia, hyperlipidaemia and hypercholesterolaemia which is  
CC associated with coronary heart disease. ABG47348-ABG59930 represent human  
CC liver single exon encoded peptides of the invention. Note: The sequence  
CC information for this patent does not appear in the printed specification  
CC but was obtained in electronic format directly from WIPO at  
CC ftp.wipo.int/pub/published\_pct\_sequences  
XX  
SQ Sequence 23 AA;  
Query Match 28.9%; Score 28; DB 4; Length 23;  
Best Local Similarity 42.9%; Pred. No. 1.3e+03;  
Matches 6; Conservative 3; Mismatches 5; Indels 0; Gaps 0;  
QY 5 LNSKIAPKIVSQEP 18  
Db | : : : |||  
4 LKSFLSIQISKQEP 17  
RESULT 220  
AAM02859  
ID AAM02859 standard; protein; 23 AA.  
XX  
AC AAM02859;  
XX  
DT 09-OCT-2001 (first entry)  
XX  
DE Peptide #1541 encoded by probe for measuring breast gene expression.  
XX  
KW Probe; human; breast disease; breast cancer; development disorder;  
KW inflammatory disease; proliferative breast disease; non-carcinoma tumour.  
XX  
OS Homo sapiens.  
XX  
PN WO200157270-A2.  
XX  
PD 09-AUG-2001.  
XX  
PF 29-JAN-2001; 2001WO-US000661.  
XX  
PR 04-FEB-2000; 2000US-0180312P.  
PR 26-MAY-2000; 2000US-0207456P.  
PR 30-JUN-2000; 2000US-00608408.  
XX

03-AUG-2000; 2000US-00632366.  
21-SEP-2000; 2000US-0234687P.  
27-SEP-2000; 2000US-0236359P.  
04-OCT-2000; 2000US-00024263.  
PR (MOLE-) MOLECULAR DYNAMICS INC.  
XX Penn SG, Hanzel DK, Chen W, Rank DR;  
XX WPI; 2002-114183/15.  
XX Spatially-addressable set of single exon nucleic acid probes, used to  
XX measure gene expression in human lung samples.  
XX Claim 27; SEQ ID NO 11599; 322pp; English.  
XX The present invention relates to novel single exon nucleic acid probes  
XX (see AA100010-AA110067). The present sequence is a peptide encoded by one  
XX such probe. The probes are useful for measuring human gene expression in  
XX a human breast sample, where the probe hybridises at high stringency to a  
XX nucleic acid expressed in the human breast. The probes are useful for  
XX predicting, diagnosing, grading, staging, monitoring and prognosing  
XX diseases of the human breast, particularly those diseases with polygenic  
XX aetiology. The diseases include: breast cancer, disorders of development,  
XX inflammatory diseases of the breast, fibrocystic changes, proliferative  
XX breast disease and non-carcinoma tumours. Note: The sequence data for  
XX this patent did not form part of the printed specification, but was  
XX obtained in electronic format directly from WIPO at  
XX ftp.wipo.int/pub/published\_pct\_sequences  
XX Sequence 23 AA;  
Query Match 28.9%; Score 28; DB 4; Length 23;  
Best Local Similarity 42.9%; Pred. No. 1.3e+03;  
Matches 6; Conservative 3; Mismatches 5; Indels 0; Gaps 0;  
QY 5 LNSKIAFKIVSQEP 18  
DB 4 LKSFLSIQISKQEP 17  
RESULT 221  
ID ABG36928 standard; peptide; 23 AA.  
XX AC ABG36928;  
XX 19-AUG-2002 (first entry)  
XX Human peptide encoded by genome-derived single exon probe SEQ ID 26593.  
XX Human; single exon probe; asthma; lung cancer; COPD; ILD;  
XX chronic obstructive pulmonary disease; interstitial lung disease;  
XX familial idiopathic pulmonary fibrosis; neurofibromatosis;  
XX tuberous sclerosis; Gaucher's disease; Niemann-Pick disease;  
XX Hermansky-Pudlak syndrome; sarcoidosis; pulmonary haemosiderosis;  
XX pulmonary histiocytosis; lymphangioleiomyomatosis; Karagener syndrome;  
XX pulmonary alveolar proteinosis; fibrocystic pulmonary dysplasia;  
XX primary ciliary dyskinesia; pulmonary hypertension;  
XX hyaline membrane disease.  
XX Homo sapiens.  
XX WO200186003-A2.  
XX 15-NOV-2001.  
XX 30-JAN-2001; 2001WO-US000665.  
XX 04-FEB-2000; 2000US-0180312P.  
XX 26-MAY-2000; 2000US-0207456P.  
XX 30-JUN-2000; 2000US-00608408.  
XX 03-AUG-2000; 2000US-00632366.  
21-SEP-2000; 2000US-0234687P.  
27-SEP-2000; 2000US-0236359P.  
04-OCT-2000; 2000GB-00024263.  
PR (MOLE-) MOLECULAR DYNAMICS INC.  
XX Penn SG, Hanzel DK, Chen W, Rank DR;  
XX WPI; 2002-114183/15.  
XX Spatially-addressable set of single exon nucleic acid probes, used to  
XX measure gene expression in human lung samples.  
XX Claim 27; SEQ ID NO 26593; 634pp; English.  
XX The invention relates to a spatially-addressable set of single exon  
XX nucleic acid probes for measuring gene expression in a sample derived  
XX from human lung comprising single exon nucleic acid probes having one of  
XX 12614 nucleic acid sequences mentioned in the specification, or their  
XX complements or the 12387 open reading frames derived from the 12614  
XX probes. Also included are a microarray comprising the novel set of probes  
XX; the novel set of probes which hybridise at high stringency to a nucleic  
XX acid expressed in the human lung; measuring gene expression in a sample  
XX derived from human lung, comprising (a) contacting the array with a  
XX collection of detectably labeled nucleic acids derived from human lung  
XX mRNA, and (b) measuring the label detectably bound to each probe of the  
XX array; identifying exons in a eukaryotic genome, comprising (a)  
XX algorithmically predicting at least one exon from genomic sequences of  
XX the eukaryote; and (b) detecting specific hybridisation of detectably  
XX labeled nucleic acids from eukaryote lung mRNA, to a single exon probe,  
XX having a fragment identical to the predicted exon, the probe is included  
XX in the above mentioned microarray; assigning exons to a single gene,  
XX comprising (a) identifying exons from genomic sequence by the method  
XX above and (b) measuring the expression of each of the exons in several  
XX tissues and/or cell types using hybridisation to a single exon  
XX microarrays having a probe with the exon, where a common pattern of  
XX expression of the exons in the tissues and/or cell types indicates that  
XX the exons should be assigned to a single gene; a peptide comprising one  
XX of 12011 sequences, mentioned in the specification, or encoded by the  
XX probes/open reading frames (ORF). The probes are used for gene expression  
XX analysis, and for identifying exons in a gene, particularly using human  
XX lung derived mRNA and for the study of lung diseases such as asthma, lung  
XX cancer, chronic obstructive pulmonary disease (COPD), interstitial lung  
XX disease (ILD), familial idiopathic pulmonary fibrosis, neurofibromatosis,  
XX tuberous sclerosis, Gaucher's disease, Niemann-Pick disease, Hermansky-  
XX Pudlak syndrome, sarcoidosis, pulmonary haemosiderosis, pulmonary  
XX histiocytosis, lymphangioleiomyomatosis, pulmonary alveolar proteinosis,  
XX Karagener syndrome, fibrocystic pulmonary dysplasia, primary ciliary  
XX dyskinesia, pulmonary hypertension and hyaline membrane disease. The  
XX present sequence is a peptide/protein encoded by a single exon probe of  
XX the invention. Note: The sequence data for this patent did not form part  
XX of the printed specification, but was obtained in electronic format  
XX directly from WIPO at ftp.wipo.int/pub/published\_pct\_sequences  
XX Sequence 23 AA;  
Query Match 28.9%; Score 28; DB 5; Length 23;  
Best Local Similarity 42.9%; Pred. No. 1.3e+03;  
Matches 6; Conservative 3; Mismatches 5; Indels 0; Gaps 0;  
QY 5 LNSKIAFKIVSQEP 18  
DB 4 LKSFLSIQISKQEP 17  
RESULT 222  
ID ADA11735 standard; protein; 23 AA.  
XX AC ADA11735;  
XX 06-NOV-2003 (first entry)  
XX





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PI Penn SG, Hanzel DK, Chen W, Rank DR;
XX WPI; 2001-488898/53.
XX Human genome-derived single exon nucleic acid probes useful for analyzing
XX gene expression in human adult liver.
XX Claim 27; SEQ ID NO 31186; 658pp; English.
XX The invention relates to a single exon nucleic acid probe (SENP) (I) for
XX measuring human gene expression in a sample derived from human adult
XX liver, comprising one of 13109 defined nucleotide sequences given in the
XX specification (or complements/ fragments). The probe hybridises at high
XX stringency to a nucleic acid molecule expressed in the human adult liver.
XX (I) may be used for predicting, measuring and displaying gene expression
XX in samples derived from human adult liver. The genes identified may be
XX involved in genetic liver diseases such as cirrhosis,
XX hyperlipoproteinaemia, hyperlipidaemia and hypercholesterolaemia which is
XX associated with coronary heart disease. ABG47348-ABG59930 represent human
XX liver single exon encoded peptides of the invention. Note: The sequence
XX information for this patent does not appear in the printed specification
XX but was obtained in electronic format directly from WIPO at
XX ftp.wipo.int/pub/published_pct_sequences
XX Sequence 25 AA;
CC Query Match 28.9%; Score 28; DB 4; Length 25;
CC Best Local Similarity 44.4%; Pred. No. 1.5e+03;
CC Matches 4; Conservative 4; Mismatches 1; Indels 0; Gaps 0;
QY 10 AFKIVSOEP 18
DB 3 AFKVMMEKP 11
|||::||
|||::||

RESULT 227
ABB09677
ID ABB09677 standard; peptide; 25 AA.
XX AC ABB09677;
XX 11-JUN-2002 (first entry)
XX Synthetic pentacosapeptide based from spinigerine.
XX Spinigerine; antibacterial; antifungal; bacterial infection;
XX fungal infection; disease resistance.
XX Synthetic.
XX WC200200836-A2.
XX 03-JAN-2002.
XX 29-JUN-2001; 2001WO-FR002100.
XX 29-JUN-2000; 2000FR-00008436.
XX (CNRS ) CNRS CENT NAT RECH SCI.
XX (ENTO-) ENTOMED.
XX Bulet P, Hoffmann J, Lamberty M;
XX WPI; 2002-171585/22.
XX New pentacosapeptides and derivatives, e.g. spinigerine isolated from
XX termices, useful as antibacterial and antifungal agents for treating or
XX preventing infections in humans, animals or plants.
XX Claim 8; Page 15; 21pp; French.
XX ABB09677-80 represent pentacosapeptides, which are based on a combination
XX of basic, hydrophobic and negatively charged or polar/large amino acid

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CC residues. The parent peptide is a natural product designated spinigerine.
CC The peptides are antibacterial and antifungal agents. The peptides may be
CC used for treating or preventing bacterial or fungal infections in humans,
CC animals or plants. They are effective against Gram positive and Gram
CC negative bacteria, filamentous fungi, yeasts and plant pathogenic
CC bacteria and fungi. Plant cells may be transformed with nucleic acid
CC sequences expressing peptides of the invention to impart disease
XX resistance to plants
XX Sequence 25 AA;
CC Query Match 28.9%; Score 28; DB 5; Length 25;
CC Best Local Similarity 36.4%; Pred. No. 1.5e+03;
CC Matches 4; Conservative 5; Mismatches 2; Indels 0; Gaps 0;
QY 4 HLNSKIAPKIV 14
DB 1 HVDKKVDKVL 11
|||::||
|||::||

RESULT 228
AAV12738
ID AAV12738 standard; protein; 26 AA.
XX AC AAV12738;
XX 21-JUN-1999 (first entry)
XX Human 5' EST secreted protein SEQ ID NO:328.
XX Human; secreted protein; EST; expressed sequence tag; diagnosis;
XX forensic; gene therapy; chromosome mapping; signal peptide;
XX upstream regulatory sequence; cytokine activity; cell proliferation;
XX differentiation; haematopoiesis regulation; tissue growth regulation;
XX reproductiv hormone regulation; chemotactic; chemokinetic; haemostatic;
XX thrombolytic; anti-inflammatory; tumour inhibition.
XX Homo sapiens.
XX WO9906549-A2.
XX 11-FEB-1999.
XX 31-JUL-1998; 98WO-IB001231.
XX 01-AUG-1997; 97US-00905279.
XX (GEST ) GENSET.
XX Dumas Milne Edwards J, Duclert A, Lacroix B;
XX WPI; 1999-153779/13.
XX N-PSDB; AAX51516.
XX New nucleic acids encoding human secreted proteins - obtained from cDNA
XX libraries derived from testis, ovary, uterus and spleen tissue.
XX Claim 34; Page 418; 522pp; English.
XX AAX51459 to AAX51691 represent 5' expressed sequence tags (ESTs) for
XX human secreted proteins, and encode the proteins given in AAY12681 to
XX AAY12913, respectively. The proteins given represent the signal peptide
XX and an N-terminal fragment of a secreted protein. The nucleic acid
XX sequences can be used for producing secreted human gene products. They
XX can also be used to develop products for diagnosis and therapy. The
XX proteins obtained may have cytokine activity, cell
XX proliferation/differentiation activity, haematopoiesis regulating
XX activity, tissue growth regulating activity, reproductiv hormone
XX regulating activity, chemotactic/ chemokinetic activity, haemostatic and
XX thrombolytic activity, receptor/ ligand activity, anti-inflammatory
XX activity, tumour inhibition activity or other activities. The products
XX can be used in forensic, gene therapy and chromosome mapping procedures.
XX The sequences can also be used for obtaining corresponding promoter

```

CC sequences. The nucleic acids encoding the signal peptide can be used for  
 CC directing extracellular secretion of a polypeptide or the insertion of a  
 CC polypeptide into a membrane, or importing a polypeptide into a cell  
 XX  
 SQ Sequence 26 AA;  
 Query Match 28.9%; Score 28; DB 2; Length 26;  
 Best Local Similarity 40.0%; Pred. No. 1.5e+03;  
 Matches 4; Conservative 4; Mismatches 2; Indels 0; Gaps 0;  
 QY 2 PNHLSKIAF 11  
 | : : : | : |  
 Db 16 PSHIDLKCSF 25  
 RESULT 229  
 ADH17225  
 ID ADH17225 standard; peptide; 27 AA.  
 XX  
 AC ADH17225;  
 XX  
 DT 11-MAR-2004 (first entry)  
 XX  
 DE Human D-AKAP2 AKB domain mutant peptide - SEQ ID 39.  
 XX  
 KW D-AKAP; dual-specific A-kinase anchor protein; regulatory subunit; PKA;  
 KW cyclic AMP-dependent protein kinase; cAMP; neurological;  
 KW neurodegenerative; Alzheimer's disease; cardiovascular; proliferative;  
 KW lipid metabolism; diabetes; obesity; retinitis pigmentosa; autoimmune;  
 KW lupus erythematosus; human; D-AKAP2; AKAP-10; SNP;  
 KW single nucleotide polymorphism; AKB domain; A-kinase binding; mutant;  
 KW mutin.  
 XX  
 OS Synthetic.  
 OS Homo sapiens.  
 FH Key Location/Qualifiers  
 FT Misc-difference 13 /note= "Wild-type Ala substituted for Asn"  
 FT Misc-difference 24 /label= Ile, Val  
 FT /note= "Residue corresponds to a SNP (single nucleotide  
 polymorphism at residue 646 within the native protein"  
 XX  
 PN WO2003093296-A2.  
 XX  
 PD 13-NOV-2003.  
 XX  
 PF 01-MAY-2003; 2003WO-US013698.  
 XX  
 PR 03-MAY-2002; 2002US-0377852P.  
 PR 07-MAR-2003; 2003US-0455408P.  
 XX  
 PA (SEQU-) SEQUENOM INC.  
 PA (REGC) UNIV CALIFORNIA.  
 XX  
 PI Braun A, Cantor CR, Kammerer SM, Taylor S, Hamuro LB, Cook C;  
 PI Olson G, Self C;  
 XX  
 WPI; 2003-903647/82.  
 DR  
 XX New isolated NOVX polypeptides and polynucleotides, useful for  
 PT preventing, diagnosing or treating NOVX-associated disorders, e.g.  
 PT osteoarthritis, obesity, atherosclerosis, cancer, Parkinson's disease,  
 PT asthma, or infections.  
 XX  
 Claim 28; SEQ ID NO 39; 149pp; English.  
 PS  
 XX The invention relates to a novel isolated mutin of a D-AKAP2 (dual-  
 CC specific A-kinase anchor protein) polypeptide where the mutin exhibits  
 CC modified binding to a regulatory subunit of PKA (cyclic AMP [cAMP]-  
 CC dependent protein kinase) compared to a native D-AKAP. The polypeptide of  
 CC the invention may be useful for treating or preventing neurological and

CC neurodegenerative disorders such as Alzheimer's disease, cardiovascular  
 CC disorders, proliferative disorders and lipid-metabolism disorders  
 CC including diabetes, obesity and retinitis pigmentosa, as well as  
 CC autoimmune disorders e.g. lupus erythematosus. The current sequence is  
 CC that of the human D-AKAP2 AKB (A-kinase binding) domain mutant peptide of  
 CC the invention.  
 XX  
 SQ Sequence 27 AA;  
 Query Match 28.9%; Score 28; DB 7; Length 27;  
 Best Local Similarity 72.7%; Pred. No. 1.6e+03;  
 Matches 8; Conservative 0; Mismatches 3; Indels 0; Gaps 0;  
 QY 5 LNSKIAPKIVS 15  
 | | | | |  
 Db 12 LNWKIAXMIVS 22  
 RESULT 230  
 ADD35429  
 ID ADD35429 standard; peptide; 28 AA.  
 XX  
 AC ADD35429;  
 XX  
 DT 15-JAN-2004 (first entry)  
 XX  
 DE Pseudomonas aeruginosa peptide deformylase peptide #1.  
 XX  
 KW crystallised recombinant protein; metabolism; Staphylococcus aureus;  
 KW Streptococcus pneumoniae; Helicobacter pylori; Escherichia coli;  
 KW Pseudomonas aeruginosa; vaccine.  
 XX  
 OS Pseudomonas aeruginosa.  
 XX  
 PN WO2003044185-A2.  
 XX  
 PD 30-MAY-2003.  
 XX  
 PF 21-NOV-2002; 2002WO-CA001769.  
 XX  
 PR 21-NOV-2001; 2001US-0332160P.  
 PR 27-NOV-2001; 2001US-0333661P.  
 PR 27-NOV-2001; 2001US-0333665P.  
 PR 18-DEC-2001; 2001US-0341770P.  
 PR 19-DEC-2001; 2001US-0341954P.  
 PR 19-DEC-2001; 2001US-0342003P.  
 PR 20-DEC-2001; 2001US-0342542P.  
 PR 21-DEC-2001; 2001US-0344252P.  
 PR 21-DEC-2001; 2001US-0343570P.  
 PR 28-DEC-2001; 2001US-0343606P.  
 PR 28-DEC-2001; 2001US-0343679P.  
 XX  
 PA (AFFI-) AFFINIUM PHARM INC.  
 XX  
 PI Edwards A, Dharamsi A, Vedadi M, Alam MZ, Awrey D, Beattie B,  
 PI Canadian V, Domagala M, Houston S, Mansoury K, Necakov S, Nethery K;  
 PI Ng I, Pinder B, Sheldrick B, Vallee F, Wrezel O;  
 XX  
 WPI; 2003-513596/48.  
 DR  
 XX New crystallized recombinant polypeptides from Staphylococcus aureus,  
 PT Streptococcus pneumoniae, Helicobacter pylori or Pseudomonas aeruginosa  
 PT involved in general metabolism, useful as drug targets for pathogenic  
 PT bacteria.  
 XX  
 PS Disclosure; SEQ ID NO 28; 277pp; English.  
 XX  
 CC The invention comprises a crystallised recombinant protein that is  
 CC involved in general metabolism, the recombinant protein may be from  
 CC Staphylococcus aureus, Streptococcus pneumoniae, Helicobacter pylori,  
 CC Escherichia coli or Pseudomonas aeruginosa. The crystallised recombinant  
 CC protein of the invention is useful in the prevention (vaccine) or  
 CC treatment of a disease or disorder caused by S. pneumoniae, H. pylori, E.

CC coli or P. aeruginosa. The present amino acid sequence was used in the  
 CC exemplification of the invention.  
 XX  
 SQ Sequence 28 AA;

Query Match 28.9%; Score 28; DB 7; Length 28;  
 Best Local Similarity 55.6%; Pred. No. 1.7e+03;  
 Matches 5; Conservative 2; Mismatches 2; Indels 0; Gaps 0;

Qy 1 EPNHLNSKI 9  
 | : ||| :  
 Db 12 ECDHLNGKL 20

RESULT 231  
 ABB43186  
 ID ABB43186 standard; peptide; 29 AA.

XX AC ABB43186;

XX DT 04-FEB-2002 (first entry)

XX DE Peptide #10692 encoded by human foetal liver single exon probe.

XX KW Human; foetal liver; gene expression; single exon nucleic acid probe.

XX OS Homo sapiens.

XX PN WO200157277-A2.

XX PD 09-AUG-2001.

XX PF 30-JAN-2001; 2001WO-US000669.

XX PR 04-FEB-2000; 2000US-0180312P.

XX PR 26-MAY-2000; 2000US-0207456P.

XX PR 30-JUN-2000; 2000US-00608408.

XX PR 03-AUG-2000; 2000US-00632366.

XX PR 21-SEP-2000; 2000US-0234687P.

XX PR 27-SEP-2000; 2000US-0236359P.

XX PR 04-OCT-2000; 2000GB-00024263.

XX PA (MOLE-) MOLECULAR DYNAMICS INC.

XX PI Penn SG, Hanzel DK, Chen W, Rank DR;

XX DR WPI; 2001-483447/52.

XX PT Human genome-derived single exon nucleic acid probes useful for analyzing

XX PS gene expression in human fetal liver.

XX PS Claim 27; SEQ ID NO 35821; 639pp + Sequence Listing; English.

XX CC The invention relates to a single exon nucleic acid probe for measuring  
 CC human gene expression in a sample derived from human foetal liver. The  
 CC single exon nucleic acid probes may be used for predicting, measuring and  
 CC displaying gene expression in samples derived from human fetal liver. The  
 CC present sequence is a peptide encoded by a single exon nucleic acid probe  
 CC of the invention. Note: the sequence data for this patent did not form  
 CC part of the printed specification, but was obtained in electronic format  
 CC directly from WIPO at ftp.wipo.int/pub/published\_pct\_sequences

XX SQ Sequence 29 AA;

Query Match 28.9%; Score 28; DB 4; Length 29;  
 Best Local Similarity 33.3%; Pred. No. 1.7e+03;  
 Matches 4; Conservative 5; Mismatches 3; Indels 0; Gaps 0;

Qy 6 NSKIAFKIVSOE 17  
 ||| : : : : :  
 Db 10 NSGISLKVIOED 21

RESULT 232

AAM37026

ID AAM37026 standard; protein; 29 AA.

XX AC AAM37026;

XX DT 17-OCT-2001 (first entry)

XX DE Peptide #11063 encoded by probe for measuring placental gene expression.

XX KW Probe; microarray; human; placenta; antenatal diagnosis;

XX KW genetic disorder.

XX OS Homo sapiens.

XX PN WO200157272-A2.

XX PD 09-AUG-2001.

XX PF 30-JAN-2001; 2001WO-US000663.

XX PR 04-FEB-2000; 2000US-0180312P.

XX PR 26-MAY-2000; 2000US-0207456P.

XX PR 30-JUN-2000; 2000US-00608408.

XX PR 03-AUG-2000; 2000US-00632366.

XX PR 21-SEP-2000; 2000US-0234687P.

XX PR 27-SEP-2000; 2000US-0236359P.

XX PR 04-OCT-2000; 2000GB-00024263.

XX PA (MOLE-) MOLECULAR DYNAMICS INC.

XX PI Penn SG, Hanzel DK, Chen W, Rank DR;

XX DR WPI; 2001-488897/53.

XX PT Human genome-derived single exon nucleic acid probes useful for analyzing

XX PS gene expression in human placenta.

XX PS Claim 27; SEQ ID NO 37295; 654pp; English.

XX CC The present invention relates to single exon nucleic acid probes (SENP;  
 CC see AAI31315-AAJ57546). The present sequence is a peptide encoded by one  
 CC such probe. The probes are useful for producing a microarray for  
 CC predicting, measuring and displaying gene expression in samples derived  
 CC from human placenta. The probes are useful for antenatal diagnosis of  
 CC human genetic disorders

XX SQ Sequence 29 AA;

Query Match 28.9%; Score 28; DB 4; Length 29;  
 Best Local Similarity 33.3%; Pred. No. 1.7e+03;  
 Matches 4; Conservative 5; Mismatches 3; Indels 0; Gaps 0;

Qy 6 NSKIAFKIVSOE 17  
 ||| : : : : :  
 Db 10 NSGISLKVIOED 21

RESULT 233

ABB26285

ID ABB26285 standard; protein; 29 AA.

XX AC ABB26285;

XX DT 23-JAN-2002 (first entry)

XX DE Protein #8284 encoded by probe for measuring heart cell gene expression.

XX KW Human; gene expression; heart; microarray; vascular system;  
 KW cardiovascular disease; hypertension; cardiac arrhythmia;  
 KW congenital heart disease.

XX OS Homo sapiens.

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XX PN WO200157274-A2.
XX XX
XX PD 09-AUG-2001.
XX XX
XX PF 30-JAN-2001; 2001WO-US000666.
XX XX
XX PR 04-FEB-2000; 2000US-0180312P.
XX PR 26-MAY-2000; 2000US-0207456P.
XX PR 30-JUN-2000; 2000US-00608408.
XX PR 03-AUG-2000; 2000US-00632366.
XX PR 21-SEP-2000; 2000US-0234687P.
XX PR 27-SEP-2000; 2000US-0236359P.
XX PR 04-OCT-2000; 2000GB-00024263.
XX XX
XX PA (MOLE-) MOLECULAR DYNAMICS INC.
XX XX
XX PI Penn SG, Hanzel DK, Chen W, Rank DR;
XX XX
XX DR WPI; 2001-488990/53.
XX XX
XX PT Single exon nucleic acid probes for analyzing gene expression in human
XX PT hearts.
XX XX
XX PS Claim 15; SEQ ID NO 28055; 530pp; English.
XX XX
XX CC The present invention relates to single exon nucleic acid probes for
XX CC measuring human gene expression in a sample derived from human heart (see
XX CC ABA21535-ABA41305). The present sequence is a protein encoded by one such
XX CC probe. The probes may be used for predicting, measuring and displaying
XX CC gene expression in samples derived from the human heart via microarrays.
XX CC By measuring gene expression, the probes are useful for predicting,
XX CC diagnosing, grading, staging, monitoring and prognosing diseases of the
XX CC human heart and vascular system e.g. cardiovascular disease,
XX CC hypertension, cardiac arrhythmias and congenital heart disease. Note: The
XX CC sequence data for this patent did not form part of the printed
XX CC specification, but was obtained in electronic format directly from WIPO
XX CC at ftp.wipo.int/pub/published_pct_sequences
XX XX
XX SQ Sequence 29 AA;
XX
Query Match 28.9%; Score 28; DB 4; Length 29;
Best Local Similarity 33.3%; Pred. No. 1.7e+03;
Matches 4; Conservative 5; Mismatches 3; Indels 0; Gaps 0;
QY 6 NSKIAPKIVSQE 17
DB |||:|:|:|:
10 NSGISLKVQIED 21
XX
RESULT 234
AMW76919
ID AAM76919 standard; protein; 29 AA.
XX AC AAM76919;
XX XX
XX DT 06-NOV-2001 (first entry)
XX XX
XX DE Human bone marrow expressed probe encoded protein SEQ ID NO: 37225.
XX XX
XX KW Human; bone marrow expressed exon; gene expression analysis; probe;
XX KW microarray; cancer; leukaemia; lymphoma; myeloma.
XX XX
XX OS Homo sapiens.
XX XX
XX PN WO200157276-A2.
XX XX
XX PD 09-AUG-2001.
XX XX
XX PF 30-JAN-2001; 2001WO-US000668.
XX XX
XX PR 04-FEB-2000; 2000US-0180312P.
XX PR 26-MAY-2000; 2000US-0207456P.

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PR 30-JUN-2000; 2000US-00608408.
PR 03-AUG-2000; 2000US-00632366.
PR 21-SEP-2000; 2000US-0234687P.
PR 27-SEP-2000; 2000US-0236359P.
PR 04-OCT-2000; 2000GB-00024263.
XX XX
XX PA (MOLE-) MOLECULAR DYNAMICS INC.
XX XX
XX PI Penn SG, Hanzel DK, Chen W, Rank DR;
XX XX
XX DR WPI; 2001-488990/53.
XX XX
XX PT Human genome-derived single exon nucleic acid probes useful for analyzing
XX PT gene expression in human bone marrow.
XX XX
XX PS Example 4; SEQ ID NO 37225; 658pp + Sequence Listing; English.
XX XX
XX CC The present invention provides a number of single exon nucleic acid
XX CC probes which are derived from genomic sequences expressed in the human
XX CC bone marrow. They can be used to measure gene expression in bone marrow
XX CC samples, which may enable the improved diagnosis and treatment of cancers
XX CC such as lymphoma, leukaemia and myeloma. The present sequence is a
XX CC protein encoded by one of the probes of the invention
XX XX
XX SQ Sequence 29 AA;
XX
Query Match 28.9%; Score 28; DB 4; Length 29;
Best Local Similarity 33.3%; Pred. No. 1.7e+03;
Matches 4; Conservative 5; Mismatches 3; Indels 0; Gaps 0;
QY 6 NSKIAPKIVSQE 17
DB |||:|:|:|:
10 NSGISLKVQIED 21
XX
RESULT 235
AMW64096
ID AAM64096 standard; protein; 29 AA.
XX AC AAM64096;
XX XX
XX DT 05-NOV-2001 (first entry)
XX XX
XX DE Human brain expressed single exon probe encoded protein SEQ ID NO: 36201.
XX XX
XX KW Human; brain expressed exon; gene expression analysis; probe; microarray;
XX KW Alzheimer's disease; multiple sclerosis; schizophrenia; epilepsy; cancer.
XX XX
XX OS Homo sapiens.
XX XX
XX PN WO200157275-A2.
XX XX
XX PD 09-AUG-2001.
XX XX
XX PF 30-JAN-2001; 2001WO-US000667.
XX XX
XX PR 04-FEB-2000; 2000US-0180312P.
XX PR 26-MAY-2000; 2000US-0207456P.
XX PR 30-JUN-2000; 2000US-00608408.
XX PR 03-AUG-2000; 2000US-00632366.
XX PR 21-SEP-2000; 2000US-0234687P.
XX PR 27-SEP-2000; 2000US-0236359P.
XX PR 04-OCT-2000; 2000GB-00024263.
XX XX
XX PA (MOLE-) MOLECULAR DYNAMICS INC.
XX XX
XX PI Penn SG, Hanzel DK, Chen W, Rank DR;
XX XX
XX DR WPI; 2001-483446/52.
XX XX
XX PT Single exon nucleic acid probes for analyzing gene expression in human
XX PT brains.
XX XX

```



PS Example 4; SEQ ID NO 36201; 650pp + Sequence Listing; English.

XX The present invention provides a number of single exon nucleic acid  
XX probes which are derived from genomic sequences expressed in the human  
XX brain. They can be used to measure gene expression in brain cell samples,  
XX which may enable the diagnosis and improved treatment of nervous system  
XX diseases such as Alzheimer's disease, multiple sclerosis, schizophrenia,  
XX epilepsy and cancers. The present sequence is a protein encoded by one of  
XX the probes of the invention

XX Sequence 29 AA;

Query Match 28.9%; Score 28; DB 4; Length 29;

Best Local Similarity 33.3%; Pred. No. 1.7e+03;

Matches 4; Conservative 5; Mismatches 3; Indels 0; Gaps 0;

QY 6 NSKIAFKIVSQE 17

DB 10 NSGISLKVIED 21

RESULT 236

ID ABG58582 standard; peptide; 29 AA.

XX ABG58582;

DT 25-FEB-2003 (first entry)

DE Human liver peptide, SEQ ID NO 37230.

XX Human; liver; cirrhosis; hyperlipoproteinaemia; hyperlipidaemia;  
XX hypercholesterolaemia; coronary heart disease.

XX Homo sapiens.

XX WO200157273-A2.

XX 09-AUG-2001.

XX 30-JAN-2001; 2001WO-US000664.

XX 04-FEB-2000; 2000US-0180312P.

XX 26-MAY-2000; 2000US-0207456P.

XX 30-JUN-2000; 2000US-00608408.

XX 03-AUG-2000; 2000US-00632366.

XX 21-SEP-2000; 2000US-0234687P.

XX 27-SEP-2000; 2000US-0236359P.

XX 04-OCT-2000; 2000GB-00024263.

XX (MOLE-) MOLECULAR DYNAMICS INC.

XX Penn SG, Hanzel DK, Chen W, Rank DR;

XX WPI; 2001-488898/53.

XX Human genome-derived single exon nucleic acid probes useful for analyzing  
XX gene expression in human adult liver.

XX Claim 27; SEQ ID NO 37230; 658pp; English.

XX The invention relates to a single exon nucleic acid probe (SENP) (I) for  
XX measuring human gene expression in a sample derived from human adult  
XX liver, comprising one of 13109 defined nucleotide sequences given in the  
XX specification (or complements/ fragments). The probe hybridises at high  
XX stringency to a nucleic acid molecule expressed in the human adult liver.  
XX (I) may be used for predicting, measuring and displaying gene expression  
XX in samples derived from human adult liver. The genes identified may be  
XX involved in genetic liver diseases such as cirrhosis.

XX hyperlipoproteinaemia, hyperlipidaemia and hypercholesterolaemia which is  
XX associated with coronary heart disease. ABG47348-ABG59930 represent human  
XX liver single exon encoded peptides of the invention. Note: The sequence  
XX information for this patent does not appear in the printed specification

CC but was obtained in electronic format directly from WIPO at  
CC ftp.wipo.int/pub/published\_pct\_sequences

XX Sequence 29 AA;

Query Match 28.9%; Score 28; DB 4; Length 29;

Best Local Similarity 33.3%; Pred. No. 1.7e+03;

Matches 4; Conservative 5; Mismatches 3; Indels 0; Gaps 0;

QY 6 NSKIAFKIVSQE 17

DB 10 NSGISLKVIED 21

RESULT 237

ABG46030

ID ABG46030 standard; peptide; 29 AA.

XX ABG46030;

DT 19-AUG-2002 (first entry)

XX Human peptide encoded by genome-derived single exon probe SEQ ID 35695.

XX Human; single exon probe; asthma; lung cancer; COPD; ILD;  
XX chronic obstructive pulmonary disease; interstitial lung disease;  
XX familial idiopathic pulmonary fibrosis; neurofibromatosis;  
XX tuberous sclerosis; Gaucher's disease; Niemann-Pick disease;  
XX Hermansky-Pudlak syndrome; sarcoidosis; pulmonary haemosiderosis;  
XX pulmonary histiocytosis; lymphangioleiomyomatosis; Karagener syndrome;  
XX pulmonary alveolar proteinosis; fibrocystic pulmonary dysplasia;  
XX primary ciliary dyskinesia; pulmonary hypertension;  
XX hyaline membrane disease.

XX Homo sapiens.

XX WO200186003-A2.

XX 15-NOV-2001.

XX 30-JAN-2001; 2001WO-US000665.

XX 04-FEB-2000; 2000US-0180312P.

XX 26-MAY-2000; 2000US-0207456P.

XX 30-JUN-2000; 2000US-00608408.

XX 03-AUG-2000; 2000US-00632366.

XX 21-SEP-2000; 2000US-0234687P.

XX 27-SEP-2000; 2000US-0236359P.

XX 04-OCT-2000; 2000GB-00024263.

XX (MOLE-) MOLECULAR DYNAMICS INC.

XX Penn SG, Hanzel DK, Chen W, Rank DR;

XX WPI; 2002-114183/15.

XX Spatially-addressable set of single exon nucleic acid probes, used to  
XX measure gene expression in human lung samples.

XX Claim 27; SEQ ID NO 35695; 634pp; English.

XX The invention relates to a spatially-addressable set of single exon  
XX nucleic acid probes for measuring gene expression in a sample derived  
XX from human lung comprising single exon nucleic acid probes having one of  
XX 12614 nucleic acid sequences mentioned in the specification, or their  
XX complements or the 12387 open reading frames derived from the 12614  
XX probes. Also included are a microarray comprising the novel set of probes  
XX; the novel set of probes which hybridise at high stringency to a nucleic  
XX acid expressed in the human lung; measuring gene expression in a sample  
XX derived from human lung, comprising (a) contacting the array with a  
XX collection of detectably labeled nucleic acids derived from human lung  
XX mRNA, and (b) measuring the label detectably bound to each probe of the  
XX array, identifying exons in a eukaryotic genome, comprising (a)

algorithmically predicting at least one exon from genomic sequences of the eukaryote; and (b) detecting specific hybridisation of detectably labeled nucleic acids from eukaryote lung mRNA, to a single exon probe, having a fragment identical to the predicted exon, the probe is included in the above mentioned microarray; assigning exons to a single gene, comprising (a) identifying exons from genomic sequence by the method above and (b) measuring the expression of each of the exons in several tissues and/or cell types using hybridisation to a single exon microarrays having a probe with the exon, where a common pattern of expression of the exons in the tissues and/or cell types indicates that the exons should be assigned to a single gene; a peptide comprising one of 12011 sequences, mentioned in the specification, or encoded by the probes/open reading frames (ORF). The probes are used for gene expression analysis, and for identifying exons in a gene, particularly using human lung derived mRNA and for the study of lung diseases such as asthma, lung cancer, chronic obstructive pulmonary disease (COPD), interstitial lung disease (ILD), familial idiopathic pulmonary fibrosis, neurofibromatosis, tuberous sclerosis, Gaucher's disease, Niemann-Pick disease, Hermansky-Pudlak syndrome, sarcoidosis, pulmonary haemosiderosis, pulmonary histiocytosis, lymphangioleiomyomatosis, pulmonary alveolar proteinosis, Karsenger syndrome, fibrocystic pulmonary dysplasia, primary ciliary dyskinesia, pulmonary hypertension and hyaline membrane disease. The present sequence is a peptide/protein encoded by a single exon probe of the invention. Note: The sequence data for this patent did not form part of the printed specification, but was obtained in electronic format directly from WIPO at [ftp.wipo.int/pub/published\\_pct\\_sequences](http://ftp.wipo.int/pub/published_pct_sequences)

CC  
XX  
SQ

Query Match 28.9%; Score 28; DB 5; Length 29;  
Best Local Similarity 33.3%; Pred. No. 1.7e+03;  
Matches 4; Conservative 5; Mismatches 3; Indels 0; Gaps 0;

QY 6 NSKIAPKIVSQE 17  
||| : : : : :  
10 NSGISLKVQED 21

RESULT 238  
AAU97881  
ID AAU97881 standard; peptide; 29 AA.  
AC AAU97881;  
DT 21-AUG-2002 (first entry)  
DE Mouse angiotensin C-terminal peptide.  
DE Plasminogen; mouse; angiotensin detection; immunological detection.  
XX Mus sp.  
XX JF2002112768-A.  
XX 16-APR-2002.  
XX 04-OCT-2000; 2000JP-00304946.  
XX 04-OCT-2000; 2000JP-00304946.  
XX (IGAK-) IGAKU SEIBUTSUGAKU KENKYUSHO KK.  
XX WPI; 2002-448751/48.  
XX Angiotensin specific binding monoclonal antibody composed of residues 79-84 of plasminogen of human being, mouse and rat used for detection of angiotensin.  
XX Example 1; Fig 2; 16pp; Japanese.  
XX The invention describes immunological detection of angiotensin with a monoclonal antibody. This sequence represents the mouse angiotensin C-terminal peptide used in the creation of an angiotensin specific binding

CC monoclonal antibody  
XX  
SQ Sequence 29 AA;  
Query Match 28.9%; Score 28; DB 5; Length 29;  
Best Local Similarity 46.2%; Pred. No. 1.7e+03;  
Matches 6; Conservative 2; Mismatches 5; Indels 0; Gaps 0;

QY 7 SKIAPKIVSQEPA 19  
||| : : : : :  
8 SVVELPTVQEPS 20

DB 8 SVVELPTVQEPS 20

RESULT 239  
AAW30507  
ID AAW30507 standard; peptide; 30 AA.  
AC AAW30507;  
DT 26-OCT-1998 (first entry)  
DE DP-1 transcription factor antagonist peptide H7.  
XX DP-1; transcription factor; antagonist; E2F protein; apoptosis;  
XX cell proliferation; cardiovascular cell; restenosis; tumour;  
KW surgical stent; therapy.  
XX Synthetic.  
OS Homo sapiens.  
XX Key Location/Qualifiers  
FT Peptide 3...9 /note= "Claim 3"  
FT Peptide 5...15 /note= "Claim 3"  
FT  
XX WO9828334-A1.  
XX 02-JUL-1998.  
XX 22-DEC-1997; 97WO-GB003506.  
XX 20-DEC-1996; 96GB-00026589.  
XX (PROL-) PROLIFIX LTD.  
XX La Thangue NB, Bandara LR;  
XX WPI; 1998-377596/32.  
XX Polypeptide fragments of the DP-1 transcription factor - used for inducing apoptosis, specifically in tumour and cardiovascular cells, e.g. for preventing re-stenosis.  
XX Claim 4; Page 44; 55pp; English.  
XX Peptide H7 comprises amino acid residues 170-199 in the DEF box (I) (see AAW30501) of transcription factor DP1. Claimed peptides (II) (see AAW30504-07) containing one or both of 2 motifs (see AAW30502-03) of the DEF box are capable of antagonising the heterodimerisation of a DP protein with an E2F protein. Also claimed are variants of these peptides, especially containing substitutions of residues corresponding to residues 167, 169, 171 and 175 of DP-1, fusion proteins (III) comprising (I) or (II) and a membrane translocation sequence (see AAW30508), expression vectors encoding (I)-(III) and host cells. (I)-(III) are used therapeutically to induce apoptosis, specifically in tumour or cardiovascular cells, either in vivo or in vitro, e.g. for purging bone marrow. Surgical stents comprising (I)-(III) are used to treat or prevent restenosis in patients who have undergone angioplasty. (I)-(III) function by inactivating the DNA-binding activity of DP/E2F heterodimers. They are also used as research reagents, as positive controls in assays for identifying antagonists of DP-1/E2F dimerisation and as immunosay agents. Also described is the use of sequences antisense to nucleic acids

CC encoding (I)-(III) to control DP levels in cells, particularly by gene  
 CC therapy. When formulated with cytotoxic or cytostatic agents, (I)-(III)  
 CC enhance cell killing

XX Sequence 30 AA;

Query Match 28.9%; Score 28; DB 2; Length 30;

Best Local Similarity 46.2%; Pred. No. 1.8e+03;

Matches 6; Conservative 3; Mismatches 4; Indels 0; Gaps 0;

QY 5 LNSKIAFKIVSQE 17

DB 4 LNVLMAMNIISKE 16

RESULT 240

AAB28613

ID AAB28613 standard; protein; 31 AA.

XX AC

XX AAB28613;

DT 12-FEB-2001 (first entry)

XX Human c-myc epitope and His6 tag peptide.

DE Human; beta-2-microglobulin; beta2M; C17;

XX haemopoietic stem/precursor cell; HSPC; mesenchymal stem cell division;

XX DNA fingerprinting; bone disease; c-myc epitope; His6.

XX Homo sapiens.

OS WO200063382-A1.

XX WO200063382-A1.

XX 26-OCT-2000.

XX 14-APR-2000; 2000WO-US009904.

XX 15-APR-1999; 99US-0129463P.

XX (OSIR-) OSIRIS THERAPEUTICS INC.

XX Liu X, Cheng L;

XX WPI; 2000-647518/62.

XX New isolated nucleic acid encoding a human hematopoietic stem/precursor

XX cell polypeptide called C17, for increasing the rate of multiplication of

XX mesenchymal stem cells in vitro and for chromosome mapping and DNA

XX fingerprinting.

XX Example 4; Page 42; 68pp; English.

XX The present sequence was used in a method for the isolation of a human

XX haematopoietic stem/precursor cell (hHSPC) polypeptide called C17. The

XX C17 polynucleotide may be used to increase the rate of multiplication of

XX human mesenchymal stem cells in vitro and to determine the presence of

XX growth-stimulating receptors on the surface of a cell. The C17

XX polypeptide may be used to generate polyclonal sera or monoclonal

XX antibodies specific for C17. The polynucleotide is used as a marker for

XX chromosome mapping and DNA fingerprinting. It is also used for detecting

XX genetic mutations in order to diagnose diseases, such as those affecting

XX bone formation

XX Sequence 31 AA;

QY Query Match 28.9%; Score 28; DB 3; Length 31;

DB Best Local Similarity 25.4%; Pred. No. 1.9e+03;

Matches 5; Conservative 7; Mismatches 5; Indels 0; Gaps 0;

QY 1 EPNHLSNKIAFKIVSQE 17

DB 2 DPSSVPSFLEOKLISEE 18

RESULT 241

AAG08759

ID AAG08759 standard; protein; 31 AA.

XX AC

XX AAG08759;

DT 17-OCT-2000 (first entry)

XX Arabidopsis thaliana protein fragment SEQ ID NO: 6423.

DE Arabidopsis thaliana.

XX Protein identification; signal transduction pathway; metabolic pathway;

XX hybridisation assay; genetic mapping; gene expression control; promoter;

XX termination sequence.

XX Arabidopsis thaliana.

OS EP1033405-A2.

XX EP1033405-A2.

XX 06-SEP-2000.

XX 25-FEB-2000; 2000EP-00301439.

XX 25-FEB-1999; 99US-0121825P.

XX 05-MAR-1999; 99US-0123180P.

XX 09-MAR-1999; 99US-0123548P.

XX 23-MAR-1999; 99US-0125788P.

XX 25-MAR-1999; 99US-0126264P.

XX 29-MAR-1999; 99US-0126785P.

XX 01-APR-1999; 99US-0127462P.

XX 06-APR-1999; 99US-0128234P.

XX 08-APR-1999; 99US-0128714P.

XX 16-APR-1999; 99US-0129845P.

XX 19-APR-1999; 99US-0130077P.

XX 21-APR-1999; 99US-0130449P.

XX 23-APR-1999; 99US-0130510P.

XX 28-APR-1999; 99US-0130891P.

XX 30-APR-1999; 99US-0131449P.

XX 30-APR-1999; 99US-0132048P.

XX 04-MAY-1999; 99US-0132407P.

XX 05-MAY-1999; 99US-0132484P.

XX 06-MAY-1999; 99US-0132485P.

XX 06-MAY-1999; 99US-0132486P.

XX 07-MAY-1999; 99US-0132487P.

XX 11-MAY-1999; 99US-0132863P.

XX 14-MAY-1999; 99US-0134256P.

XX 14-MAY-1999; 99US-0134218P.

XX 14-MAY-1999; 99US-0134219P.

XX 14-MAY-1999; 99US-0134221P.

XX 14-MAY-1999; 99US-0134370P.

XX 18-MAY-1999; 99US-0134768P.

XX 19-MAY-1999; 99US-0134941P.

XX 20-MAY-1999; 99US-0135124P.

XX 21-MAY-1999; 99US-0135353P.

XX 24-MAY-1999; 99US-0135629P.

XX 25-MAY-1999; 99US-0136021P.

XX 27-MAY-1999; 99US-0136392P.

XX 28-MAY-1999; 99US-0136782P.

XX 03-JUN-1999; 99US-0137222P.

XX 03-JUN-1999; 99US-0137528P.

XX 04-JUN-1999; 99US-0137502P.

XX 07-JUN-1999; 99US-0137724P.

XX 08-JUN-1999; 99US-0138094P.

XX 10-JUN-1999; 99US-0138540P.

XX 14-JUN-1999; 99US-0138847P.

XX 14-JUN-1999; 99US-0139119P.

XX 16-JUN-1999; 99US-0139452P.

XX 16-JUN-1999; 99US-0139453P.

XX 17-JUN-1999; 99US-0139492P.

XX 18-JUN-1999; 99US-0139454P.

XX 18-JUN-1999; 99US-0139455P.

XX 18-JUN-1999; 99US-0139456P.

XX 18-JUN-1999; 99US-0139457P.

PR 18-JUN-1999; 99US-0139458P.  
PR 18-JUN-1999; 99US-0139459P.  
PR 18-JUN-1999; 99US-0139460P.  
PR 18-JUN-1999; 99US-0139461P.  
PR 18-JUN-1999; 99US-0139462P.  
PR 18-JUN-1999; 99US-0139463P.  
PR 18-JUN-1999; 99US-0139750P.  
PR 18-JUN-1999; 99US-0139763P.  
PR 21-JUN-1999; 99US-0139817P.  
PR 22-JUN-1999; 99US-0139899P.  
PR 23-JUN-1999; 99US-0140353P.  
PR 23-JUN-1999; 99US-0140354P.  
PR 24-JUN-1999; 99US-0140696P.  
PR 28-JUN-1999; 99US-0140823P.  
PR 29-JUN-1999; 99US-0140991P.  
PR 30-JUN-1999; 99US-0141287P.  
PR 01-JUL-1999; 99US-0141842P.  
PR 01-JUL-1999; 99US-0142154P.  
PR 02-JUL-1999; 99US-0142055P.  
PR 06-JUL-1999; 99US-0142390P.  
PR 08-JUL-1999; 99US-0142803P.  
PR 09-JUL-1999; 99US-0142920P.  
PR 12-JUL-1999; 99US-0142977P.  
PR 13-JUL-1999; 99US-0143542P.  
PR 14-JUL-1999; 99US-0143624P.  
PR 15-JUL-1999; 99US-0144005P.  
PR 16-JUL-1999; 99US-0144085P.  
PR 16-JUL-1999; 99US-0144086P.  
PR 19-JUL-1999; 99US-0144325P.  
PR 19-JUL-1999; 99US-0144331P.  
PR 19-JUL-1999; 99US-0144332P.  
PR 19-JUL-1999; 99US-0144333P.  
PR 19-JUL-1999; 99US-0144334P.  
PR 20-JUL-1999; 99US-0144335P.  
PR 20-JUL-1999; 99US-0144632P.  
PR 20-JUL-1999; 99US-0144684P.  
PR 21-JUL-1999; 99US-0144814P.  
PR 21-JUL-1999; 99US-0145086P.  
PR 21-JUL-1999; 99US-0145088P.  
PR 22-JUL-1999; 99US-0145085P.  
PR 22-JUL-1999; 99US-0145087P.  
PR 22-JUL-1999; 99US-0145089P.  
PR 22-JUL-1999; 99US-0145192P.  
PR 23-JUL-1999; 99US-0145145P.  
PR 23-JUL-1999; 99US-0145218P.  
PR 23-JUL-1999; 99US-0145224P.  
PR 26-JUL-1999; 99US-0145276P.  
PR 27-JUL-1999; 99US-0145913P.  
PR 27-JUL-1999; 99US-0145918P.  
PR 28-JUL-1999; 99US-0145919P.  
PR 28-JUL-1999; 99US-0145951P.  
PR 02-AUG-1999; 99US-0146386P.  
PR 02-AUG-1999; 99US-0146388P.  
PR 03-AUG-1999; 99US-0147038P.  
PR 04-AUG-1999; 99US-0147204P.  
PR 04-AUG-1999; 99US-0147302P.  
PR 05-AUG-1999; 99US-0147192P.  
PR 05-AUG-1999; 99US-0147260P.  
PR 06-AUG-1999; 99US-0147303P.  
PR 09-AUG-1999; 99US-0147416P.  
PR 09-AUG-1999; 99US-0147493P.  
PR 09-AUG-1999; 99US-0147935P.  
PR 10-AUG-1999; 99US-0148171P.  
PR 11-AUG-1999; 99US-0148319P.  
PR 12-AUG-1999; 99US-0148341P.  
PR 13-AUG-1999; 99US-0148565P.  
PR 13-AUG-1999; 99US-0148684P.  
PR 16-AUG-1999; 99US-0149368P.  
PR 17-AUG-1999; 99US-0149175P.  
PR 18-AUG-1999; 99US-0149426P.  
PR 20-AUG-1999; 99US-0149722P.

PR 20-AUG-1999; 99US-0149723P.  
PR 20-AUG-1999; 99US-0149923P.  
PR 23-AUG-1999; 99US-0149902P.  
PR 23-AUG-1999; 99US-0149930P.  
PR 25-AUG-1999; 99US-0150566P.  
PR 26-AUG-1999; 99US-0150884P.  
PR 27-AUG-1999; 99US-0151065P.  
PR 27-AUG-1999; 99US-0151066P.  
PR 27-AUG-1999; 99US-0151080P.  
PR 30-AUG-1999; 99US-0151303P.  
PR 31-AUG-1999; 99US-0151438P.  
PR 01-SEP-1999; 99US-0151930P.  
PR 07-SEP-1999; 99US-0152363P.  
PR 10-SEP-1999; 99US-0153070P.  
PR 13-SEP-1999; 99US-0153758P.  
PR 15-SEP-1999; 99US-0154018P.  
PR 16-SEP-1999; 99US-0154039P.  
PR 20-SEP-1999; 99US-0154779P.  
PR 22-SEP-1999; 99US-0155139P.  
PR 23-SEP-1999; 99US-0155486P.  
PR 24-SEP-1999; 99US-0155659P.  
PR 28-SEP-1999; 99US-0156458P.  
PR 29-SEP-1999; 99US-0156596P.  
PR 04-OCT-1999; 99US-0157117P.  
PR 05-OCT-1999; 99US-0157753P.  
PR 06-OCT-1999; 99US-0157865P.  
PR 07-OCT-1999; 99US-0158029P.  
PR 08-OCT-1999; 99US-0158232P.  
PR 12-OCT-1999; 99US-0158369P.  
PR 13-OCT-1999; 99US-0159293P.  
PR 13-OCT-1999; 99US-0159294P.  
PR 13-OCT-1999; 99US-0159295P.  
PR 14-OCT-1999; 99US-0159329P.  
PR 14-OCT-1999; 99US-0159330P.  
PR 14-OCT-1999; 99US-0159331P.  
PR 14-OCT-1999; 99US-0159637P.  
PR 14-OCT-1999; 99US-0159638P.  
PR 18-OCT-1999; 99US-0159584P.  
PR 21-OCT-1999; 99US-0160741P.  
PR 21-OCT-1999; 99US-0160767P.  
PR 21-OCT-1999; 99US-0160768P.  
PR 21-OCT-1999; 99US-0160770P.  
PR 21-OCT-1999; 99US-0160814P.  
PR 21-OCT-1999; 99US-0160815P.  
PR 22-OCT-1999; 99US-0160980P.  
PR 22-OCT-1999; 99US-0160981P.  
PR 22-OCT-1999; 99US-0160989P.  
PR 25-OCT-1999; 99US-0161404P.  
PR 25-OCT-1999; 99US-0161405P.  
PR 25-OCT-1999; 99US-0161406P.  
PR 26-OCT-1999; 99US-0161359P.  
PR 26-OCT-1999; 99US-0161360P.  
PR 26-OCT-1999; 99US-0161361P.  
PR 28-OCT-1999; 99US-0161920P.  
PR 28-OCT-1999; 99US-0161992P.  
PR 28-OCT-1999; 99US-0161993P.  
PR 29-OCT-1999; 99US-0162142P.

Query Match 28.9%; Score 28; DB 3; Length 31;  
Best Local Similarity 35.7%; Pred. No. 1.9e+03;  
Matches 5; Conservative 5; Mismatches 4; Indels 0; Gaps 0;

QY ||| : : : : :  
Db 4 NSPLSKALTRGPA 17

RESULT 242  
ABU02095  
ID ABU02095 standard; protein; 34 AA.  
XX AC ABU02095;  
XX



AC AAR58093;  
 XX  
 DT 20-SEP-1994 (first entry)  
 XX  
 DE [Asn13]-hPTH(1-38)-OH.  
 XX Human parathyroid hormone; hPTH; variant; analogue; calcium; depletion;  
 KW fixation; resorption; osteopathy; osteoporosis; hypoparathyroidism.  
 XX  
 OS Synthetic.  
 XX  
 PN GB2269176-A.  
 XX  
 PD 02-FEB-1994.  
 XX  
 XX 12-JUL-1993; 93GB-00014384.  
 PF  
 XX 15-JUL-1992; 92GB-00015009.  
 PR  
 PR 18-DEC-1992; 92GB-00026415.  
 PR  
 PR 23-DEC-1992; 92GB-00026859.  
 PR  
 PR 23-DEC-1992; 92GB-00026861.  
 PR  
 PR 28-JAN-1993; 93GB-00001691.  
 PR  
 PR 28-JAN-1993; 93GB-00001692.  
 PR  
 PR 14-APR-1993; 93GB-00007673.  
 PR  
 PR 19-APR-1993; 93GB-00008033.  
 XX  
 PA (SANO ) SANDOZ LTD.  
 XX  
 XX Lewis I, Schneider H, Waelchli R, Rainer A;  
 PI  
 XX  
 DR WPI; 1994-018352/03.  
 XX  
 XX New active para-thyroid hormone variants - used for treating or  
 PT preventing osteoporosis etc.  
 PT  
 XX Example 90; Page 38; 92pp; English.  
 PS  
 XX This peptide is an example of a highly generic formula covering  
 CC parathyroid hormone variants useful for treating or preventing bone  
 CC conditions associated with calcium depletion/resorption, in cases where  
 CC calcium fixation is required (esp. osteoporosis) or to treat  
 CC hypoparathyroidism  
 XX  
 XX Sequence 38 AA;  
 SQ  
 Query Match 28.9%; Score 28; DB 2; Length 38;  
 Best Local Similarity 100.0%; Pred. No. 2.4e+03;  
 Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
 QY 3 NHLNS 7  
 DB 13 NHLNS 17  
 RESULT 245  
 AAY13001  
 ID AAY13001 standard; protein; 38 AA.  
 XX  
 AC AAY13001;  
 XX  
 XX 22-JUN-1999 (first entry)  
 DT  
 XX Human secreted protein encoded by 5' EST SEQ ID NO: 15.  
 DE  
 XX Human; secreted protein; EST; expressed sequence tag; diagnosis;  
 KW forensic; gene therapy; chromosome mapping; signal peptide;  
 KW upstream regulatory sequence; cytokine activity; cell proliferation;  
 KW differentiation; haematopoiesis regulation; tissue growth regulation;  
 KW reproductively hormone regulation; chemotactic; chemokinetic; haemostatic;  
 KW thrombolytic; anti-inflammatory; tumour inhibition.  
 XX  
 OS Homo sapiens.  
 XX

PN WO9906552-A2.  
 XX  
 PD 11-FEB-1999.  
 XX  
 PF 31-JUL-1998; 98WO-IB001236.  
 XX  
 PR 01-AUG-1997; 97US-00905223.  
 XX  
 PA (GEST ) GENSET.  
 XX  
 XX Dumas Milne Edwards J, Duclert A, Lacroix B;  
 PI  
 XX WPI; 1999-153782/13.  
 DR  
 DR N-PSDB; AAX51801.  
 XX  
 PT New isolated brain-derived nucleic acids - used to develop products which  
 PT may have cytokine, immune, regulatory, haematopoiesis regulating, anti-  
 PT inflammatory or tumour inhibition activity.  
 XX  
 PS Claim 34; Page 444; 577pp; English.  
 XX  
 CC AAX51787 to AAX52019 represent 5' expressed sequence tags (ESTs) for  
 CC human secreted proteins, and encode the proteins given in AAY12987 to  
 CC AAY13219, respectively. The proteins given represent the signal peptide  
 CC and an N-terminal fragment of a secreted protein. The nucleic acid  
 CC sequences can be used for producing secreted human gene products. They  
 CC can also be used to develop products for diagnosis and therapy. The  
 CC proteins obtained may have cytokine activity, cell  
 CC proliferation/differentiation activity, haematopoiesis regulating  
 CC activity, tissue growth regulating activity, reproductive hormone  
 CC regulating activity, chemotactic/ chemokinetic activity, haemostatic and  
 CC thrombolytic activity, receptor/ ligand activity, anti-inflammatory  
 CC activity, tumour inhibition activity or other activities. The products  
 CC can be used in forensic, gene therapy and chromosome mapping procedures.  
 CC The sequences can also be used for obtaining corresponding promoter  
 CC sequences. The nucleic acids encoding the signal peptide can be used for  
 CC directing extracellular secretion of a polypeptide or the insertion of a  
 CC polypeptide into a membrane, or importing a polypeptide into a cell  
 XX  
 XX Sequence 38 AA;  
 SQ  
 Query Match 28.9%; Score 28; DB 2; Length 38;  
 Best Local Similarity 66.7%; Pred. No. 2.4e+03;  
 Matches 4; Conservative 2; Mismatches 0; Indels 0; Gaps 0;  
 QY 2 NHLNS 7  
 DB 24 PSHNS 29  
 RESULT 246  
 AAM15186  
 ID AAM15186 standard; protein; 38 AA.  
 XX  
 AC AAM15186;  
 XX  
 DT 12-OCT-2001 (first entry)  
 XX  
 DE Peptide #1620 encoded by probe for measuring cervical gene expression.  
 XX  
 KW Probe; human; microarray; gene expression; cervical epithelial cell;  
 KW cervical cancer.  
 XX  
 OS Homo sapiens.  
 XX  
 XX WO200157278-A2.  
 PN  
 PD 09-AUG-2001.  
 XX  
 PF 30-JAN-2001; 2001WO-US000670.  
 XX  
 XX 04-FEB-2000; 2000US-0180312P.  
 PR  
 PR 26-MAY-2000; 2000US-0207456P.

PR 30-JUN-2000; 2000US-00608408.  
PR 03-AUG-2000; 2000US-00632366.  
PR 21-SEP-2000; 2000US-0234687P.  
PR 27-SEP-2000; 2000US-0236359P.  
PR 04-OCT-2000; 2000GB-00024263.  
XX  
XX (MOLE-) MOLECULAR DYNAMICS INC.  
XX PA  
XX Penn SG, Hanzel DK, Chen W, Rank DR;  
XX WPI; 2001-488901/53.  
XX  
XX Human genome-derived single exon nucleic acid probes useful for analyzing  
PT gene expression in human cervical epithelial cells.  
XX  
XX Claim 27; SEQ ID NO 20012; 487pp; English.  
XX  
XX The present invention relates to human single exon nucleic acid probes  
CC (SENP: see AAI10068-AAI28459). The present sequence is a peptide encoded  
CC by one such probe. The SENPs are derived from human Hela cells. The SENPs  
CC can be used to produce a single exon microarray, which can be used for  
CC measuring human gene expression in a sample derived from human cervical  
CC epithelial cells. By measuring gene expression, the probes are therefore  
CC useful in grading and/or staging of diseases of the cervix, notably  
CC cervical cancer. Note: The sequence data for this patent did not form  
CC part of the printed specification, but was obtained in electronic format  
CC directly from WIPO at ftp.wipo.int/pub/published\_pct\_sequences  
XX  
XX Sequence 38 AA;  
SQ

Query Match 28.9%; Score 28; DB 4; Length 38;  
Best Local Similarity 50.0%; Pred. No. 2.4e+03;  
Matches 5; Conservative 2; Mismatches 3; Indels 0; Gaps 0;  
QY 2 PNHLNSKIAP 11  
||| | : :  
Db 23 PNHDNKQOSF 32

RESULT 247  
ABB34179  
ID ABB34179 standard; peptide; 38 AA.  
XX AC  
XX ABB34179;  
XX  
XX 04-FEB-2002 (first entry)  
XX  
XX Peptide #1685 encoded by human foetal liver single exon probe.  
XX Human; foetal liver; gene expression; single exon nucleic acid probe.  
XX Homo sapiens.  
XX  
XX WO200157277-A2.  
XX  
XX 09-AUG-2001.  
XX  
XX 30-JAN-2001; 2001WO-US000669.  
XX  
XX 04-FEB-2000; 2000US-0180312P.  
XX 26-MAY-2000; 2000US-0207456P.  
XX 30-JUN-2000; 2000US-00608408.  
XX 03-AUG-2000; 2000US-00632366.  
XX 21-SEP-2000; 2000US-0234687P.  
XX 27-SEP-2000; 2000US-0236359P.  
XX 04-OCT-2000; 2000GB-00024263.  
XX  
XX (MOLE-) MOLECULAR DYNAMICS INC.  
XX PA  
XX Penn SG, Hanzel DK, Chen W, Rank DR;  
XX WPI; 2001-483447/52.  
XX

PT Human genome-derived single exon nucleic acid probes useful for analyzing  
PT gene expression in human foetal liver.  
XX  
XX Claim 27; SEQ ID NO 26814; 639pp + Sequence Listing; English.  
XX  
XX The invention relates to a single exon nucleic acid probe for measuring  
CC human gene expression in a sample derived from human foetal liver. The  
CC single exon nucleic acid probes may be used for predicting, measuring and  
CC displaying gene expression in samples derived from human foetal liver. The  
CC present sequence is a peptide encoded by a single exon nucleic acid probe  
CC of the invention. Note: The sequence data for this patent did not form  
CC part of the printed specification, but was obtained in electronic format  
CC directly from WIPO at ftp.wipo.int/pub/published\_pct\_sequences  
XX  
XX Sequence 38 AA;  
SQ

Query Match 28.9%; Score 28; DB 4; Length 38;  
Best Local Similarity 50.0%; Pred. No. 2.4e+03;  
Matches 5; Conservative 2; Mismatches 3; Indels 0; Gaps 0;  
QY 2 PNHLNSKIAP 11  
||| | : :  
Db 23 PNHDNKQOSF 32

RESULT 248  
AAM27645  
ID AAM27645 standard; protein; 38 AA.  
XX AC  
XX AAM27645;  
XX  
XX 17-OCT-2001 (first entry)  
XX  
XX Peptide #1682 encoded by probe for measuring placental gene expression.  
XX Probe; microarray; human; placenta; antenatal diagnosis;  
XX genetic disorder.  
XX Homo sapiens.  
XX  
XX WO200157272-A2.  
XX  
XX 09-AUG-2001.  
XX  
XX 30-JAN-2001; 2001WO-US000663.  
XX  
XX 04-FEB-2000; 2000US-0180312P.  
XX 26-MAY-2000; 2000US-0207456P.  
XX 30-JUN-2000; 2000US-00608408.  
XX 03-AUG-2000; 2000US-00632366.  
XX 21-SEP-2000; 2000US-0234687P.  
XX 27-SEP-2000; 2000US-0236359P.  
XX 04-OCT-2000; 2000GB-00024263.  
XX  
XX (MOLE-) MOLECULAR DYNAMICS INC.  
XX PA  
XX Penn SG, Hanzel DK, Chen W, Rank DR;  
XX WPI; 2001-488897/53.  
XX  
XX Human genome-derived single exon nucleic acid probes useful for analyzing  
PT gene expression in human placenta.  
XX  
XX Claim 27; SEQ ID NO 27914; 654pp; English.  
XX  
XX The present invention relates to single exon nucleic acid probes (SENP:  
CC see AAI31315-AAI57546). The present sequence is a peptide encoded by one  
CC such probe. The probes are useful for producing a microarray for  
CC predicting, measuring and displaying gene expression in samples derived  
CC from human placenta. The probes are useful for antenatal diagnosis of  
CC human genetic disorders  
XX  
XX Sequence 38 AA;  
SQ

Query Match 28.9%; Score 28; DB 4; Length 38;  
Best Local Similarity 50.0%; Pred. No. 2.4e+03;  
Matches 5; Conservative 2; Mismatches 3; Indels 0; Gaps 0;

QY 2 PNHLNSKIAP 11  
||| | : :  
Db 23 PNHDNKQOSF 32

RESULT 249  
ABB29012  
ID ABB29012 standard; peptide; 38 AA.  
XX  
AC ABB29012;  
XX  
DT 01-FEB-2002 (first entry)  
XX  
DE Peptide #1663 encoded by breast cell single exon nucleic acid probe.  
XX  
KW Human; microarray; single exon probe; gene expression; breast; disease;  
XX cancer.  
XX  
OS Homo sapiens.  
XX  
PN WO200157271-A2.  
XX  
PD 09-AUG-2001.  
XX  
PF 30-JAN-2001; 2001WO-US000662.  
XX  
PR 04-FEB-2000; 2000US-0180312P.  
PR 26-MAY-2000; 2000US-0207456P.  
PR 30-JUN-2000; 2000US-00608408.  
PR 03-AUG-2000; 2000US-00632366.  
PR 21-SEP-2000; 2000US-0234687P.  
PR 27-SEP-2000; 2000US-0236359P.  
PR 04-OCT-2000; 2000GB-00024263.  
XX  
PA (MOLE-) MOLECULAR DYNAMICS INC.

Penn SG, Hanzel DK, Chen W, Rank DR;

WPI; 2001-496933/54.

New spatially-addressable set of single exon nucleic acid probes, useful  
for measuring gene expression in sample derived from human breast,  
comprises number of single exon nucleic acid probes.

Claim 27; SEQ ID NO 11980; 327pp + Sequence Listing; English.

The invention relates to a spatially-addressable set of single exon  
nucleic acid probes for measuring gene expression in a sample derived  
from human breast and BT 474 cells. The method involves contacting the  
probes with a collection of detectably labelled nucleic acids derived  
from mRNA of human breast, and then measuring the label bound to each  
probe of the microarray. The probes are useful for verifying the  
expression of regions of genomic DNA predicted to encode proteins. They  
are useful for gene discovery, and for determining predisposition and/or  
prognosing breast disease. Gene expression analysis is useful for  
assessing the toxicity of chemical agents on cells. The microarray of  
this invention presents a far greater diversity of probes for measuring  
gene expression, with far less bias than expressed sequence tag  
microarrays. The method is suitable for rapid production of functional  
information from genomic sequence. The present sequence is a peptide  
encoded by a single exon nucleic acid probe of the invention. Note: The  
sequence data for this patent did not form part of the printed  
specification, but was obtained in electronic format directly from WIPO  
at ftp.wipo.int/pub/published\_pct\_sequences

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PR 26-MAY-2000; 2000US-0207456P.  
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PR 04-OCT-2000; 2000GB-00024263.  
XX  
PA (MOLE-) MOLECULAR DYNAMICS INC.

Penn SG, Hanzel DK, Chen W, Rank DR;

WPI; 2001-488899/53.

Single exon nucleic acid probes for analyzing gene expression in human  
hearts.

Claim 15; SEQ ID NO 21390; 530pp; English.

The present invention relates to single exon nucleic acid probes for  
measuring human gene expression in a sample derived from human heart (see  
ABA21535-ABA41305). The present sequence is a protein encoded by one such  
probe. The probes may be used for predicting, measuring and displaying  
gene expression in samples derived from the human heart via microarrays.  
By measuring gene expression, the probes are useful for predicting, the  
diagnosing, grading, staging, monitoring and prognosing diseases of the  
human heart and vascular system e.g. cardiovascular disease,  
hypertension, cardiac arrhythmias and congenital heart disease. Note: The  
sequence data for this patent did not form part of the printed  
specification, but was obtained in electronic format directly from WIPO  
at ftp.wipo.int/pub/published\_pct\_sequences

Sequence 38 AA;

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Best Local Similarity 50.0%; Pred. No. 2.4e+03;  
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## ALIGNMENTS

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; Publication No. US20030216544A1
; GENERAL INFORMATION:
; APPLICANT: Harley, John
; TITLE OF INVENTION: METHODS AND REAGENTS FOR DIAGNOSIS OF
; AUTOCITIBODIES
; NUMBER OF SEQUENCES: 218
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Patrea L. Pabst
; STREET: Suite 2000, 1201 West Peachtree Street, N.E.
; CITY: Atlanta
; STATE: GA
; COUNTRY: USA
; ZIP: 30309-3400
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
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SOFTWARE: PatentIn Release #1.0, Version #1.25  
CURRENT APPLICATION DATA:  
APPLICATION NUMBER: US/10/376,121A  
FILING DATE: 27-Mar-2003  
CLASSIFICATION: <Unknown>  
PRIOR APPLICATION DATA:  
APPLICATION NUMBER: 07/867,819  
FILING DATE: April 13, 1992  
APPLICATION NUMBER: 07/648,205  
FILING DATE: January 31, 1991  
APPLICATION NUMBER: 07/472,947  
FILING DATE: January 31, 1990  
ATTORNEY/AGENT INFORMATION:  
NAME: Pabst, Patrea L.  
REGISTRATION NUMBER: 31,284  
REFERENCE/DOCKET NUMBER: OMRfil4CIP(2)DIV(2)  
TELECOMMUNICATION INFORMATION:  
TELEPHONE: (404)-817-8473  
TELEFAX: (404)-817-8588  
INFORMATION FOR SEQ ID NO: 55:  
SEQUENCE CHARACTERISTICS:  
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TYPE: amino acid  
STRANDEDNESS: single  
TOPOLOGY: linear  
MOLECULE TYPE: peptide  
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; Publication No. US20040031072A1  
; GENERAL INFORMATION:  
; APPLICANT: La Rosa Thomas J  
; APPLICANT: Kovalic David K  
; APPLICANT: Zhou Yihua  
; APPLICANT: Cao Yongwei  
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With  
; FILE REFERENCE: 38-21(53223)B  
; CURRENT APPLICATION NUMBER: US/10/424,599  
; CURRENT FILING DATE: 2003-04-28  
; NUMBER OF SEQ ID NOS: 285684  
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; Publication No. US20040031072A1  
; GENERAL INFORMATION:  
; APPLICANT: La Rosa Thomas J  
; APPLICANT: Kovalic David K  
; APPLICANT: Zhou Yihua  
; APPLICANT: Cao Yongwei  
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With  
; FILE REFERENCE: 38-21(53223)B  
; CURRENT APPLICATION NUMBER: US/10/424,599  
; CURRENT FILING DATE: 2003-04-28  
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; APPLICANT: La Rosa Thomas J  
; APPLICANT: Kovalic David K  
; APPLICANT: Zhou Yihua  
; APPLICANT: Cao Yongwei  
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With  
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; OTHER INFORMATION: Clone ID: PAT\_MRT3847\_98372C.1.pep  
US-10-424-599-283885

Query Match 37.1%; Score 36; DB 15; Length 50;  
Best Local Similarity 41.2%; Pred. No. 1.5e+02;  
Matches 7; Conservative 2; Mismatches 8; Indels 0; Gaps 0;

QY 2 PNLNSKIAPKIVSQEP 18  
DB 21 PAHLNKKNCFLINWP 37

## RESULT 5

US-10-424-599-223392  
; Sequence 223392, Application US/10424599  
; Publication No. US20040031072A1  
; GENERAL INFORMATION:  
; APPLICANT: La Rosa Thomas J  
; APPLICANT: Kovalic David K  
; APPLICANT: Zhou Yihua  
; APPLICANT: Cao Yongwei

Query Match	35.1%;	Score 34;	DB 15;	Length 38;
Best Local Similarity	61.5%;	Pred. No. 2.4e+02;		
Matches	8.	Conservative	1;	Mismatches 4;
				Indels 0;
				Gaps 0;

;	TITLE OF INVENTION:	HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES USEFUL FOR
:	TITLE OF INVENTION:	GENE EXPRESSION ANALYSIS BY MICROARRAY



;; PRIOR FILING DATE: 2000-09-27  
;; PRIOR APPLICATION NUMBER: PCT/US01/00666  
;; PRIOR FILING DATE: 2001-01-30  
;; PRIOR APPLICATION NUMBER: PCT/US01/00667  
;; PRIOR FILING DATE: 2001-01-30  
;; PRIOR APPLICATION NUMBER: PCT/US01/00664  
;; PRIOR FILING DATE: 2001-01-30  
;; PRIOR APPLICATION NUMBER: PCT/US01/00669  
;; PRIOR FILING DATE: 2001-01-30  
;; PRIOR APPLICATION NUMBER: PCT/US01/00665  
;; PRIOR FILING DATE: 2001-01-30  
;; PRIOR APPLICATION NUMBER: PCT/US01/00668  
;; PRIOR FILING DATE: 2001-01-30  
;; PRIOR APPLICATION NUMBER: PCT/US01/00663  
;; PRIOR FILING DATE: 2001-01-30  
;; PRIOR APPLICATION NUMBER: PCT/US01/00662  
;; PRIOR FILING DATE: 2001-01-30  
;; PRIOR APPLICATION NUMBER: PCT/US01/00661  
;; PRIOR FILING DATE: 2001-01-30  
;; PRIOR APPLICATION NUMBER: PCT/US01/00670  
;; PRIOR FILING DATE: 2001-01-30  
;; PRIOR APPLICATION NUMBER: US 60/234,687  
;; PRIOR FILING DATE: 2000-09-21  
;; PRIOR APPLICATION NUMBER: US 09/608,408  
;; PRIOR FILING DATE: 2000-06-30  
;; PRIOR APPLICATION NUMBER: US 09/774,203  
;; PRIOR FILING DATE: 2001-01-29  
;; NUMBER OF SEQ ID NOS: 49117  
;; SOFTWARE: Anomax Sequence Listing Engine vers. 1.1  
;; SEQ ID NO 43939  
;; LENGTH: 26  
;; TYPE: PRT  
;; ORGANISM: Homo sapiens  
;; FEATURE:  
;; OTHER INFORMATION: MAP TO AC004827.1  
;; OTHER INFORMATION: EXPRESSED IN ADULT LIVER, SIGNAL = 0.56  
;; OTHER INFORMATION: EXPRESSED IN PLACENTA, SIGNAL = 0.59  
;; OTHER INFORMATION: EXPRESSED IN LUNG, SIGNAL = 0.64  
;; OTHER INFORMATION: EXPRESSED IN BRAIN, SIGNAL = 0.71  
;; OTHER INFORMATION: EXPRESSED IN BONE MARROW, SIGNAL = 0.7  
US-09-864-761-43939

Query Match 34.0%; Score 33; DB 9; Length 26;  
Best Local Similarity 43.8%; Pred. No. 2.3e+02;  
Matches 7; Conservative 2; Mismatches 7; Indels 0; Gaps 0;

Qy 3 NHLNFKIAFKIVSQEP 18  
Db 2 NTLKRTPIQLGQEP 17

RESULT 10  
US-10-424-599-144376  
; Sequence 144376, Application US/10424599  
; Publication No. US20040031072A1  
; GENERAL INFORMATION:  
; APPLICANT: La Rosa Thomas J  
; APPLICANT: Kovalic David K  
; APPLICANT: Zhou Yihua  
; APPLICANT: Cao Yongwei  
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With  
; FILE REFERENCE: 38-21(53223)B  
; CURRENT APPLICATION NUMBER: US/10/424,599  
; CURRENT FILING DATE: 2003-04-28  
; NUMBER OF SEQ ID NOS: 285684  
; SEQ ID NO 144376  
; LENGTH: 35  
; TYPE: PRT  
; ORGANISM: Glycine max  
; FEATURE:  
; OTHER INFORMATION: Clone ID: PAT\_MRT3847\_101385C.1.pap  
US-10-424-599-144376

Query Match 34.0%; Score 33; DB 15; Length 35;  
Best Local Similarity 45.5%; Pred. No. 3.3e+02;  
Matches 5; Conservative 4; Mismatches 2; Indels 0; Gaps 0;

Qy 3 NHLNFKIAFKI 13  
Db 15 NHNTLLHYKI 25

RESULT 11  
US-10-437-963-164005  
; Sequence 164005, Application US/10437963  
; Publication No. US20040123343A1  
; GENERAL INFORMATION:  
; APPLICANT: La Rosa, Thomas J.  
; APPLICANT: Kovalic, David K.  
; APPLICANT: Zhou, Yihua  
; APPLICANT: Cao, Yongwei  
; APPLICANT: Wu, Wei  
; APPLICANT: Boukharov, Andrey A.  
; APPLICANT: Barbazuk, Brad  
; APPLICANT: Li, Ping  
; TITLE OF INVENTION: Rice Nucleic Acid Molecules and Other Molecules Associated With  
; FILE REFERENCE: 38-21(53221)B  
; CURRENT APPLICATION NUMBER: US/10/437,963  
; CURRENT FILING DATE: 2003-05-14  
; NUMBER OF SEQ ID NOS: 204966  
; SEQ ID NO 164005  
; LENGTH: 38  
; TYPE: PRT  
; ORGANISM: Oryza sativa  
; FEATURE:  
; OTHER INFORMATION: Clone ID: PAT\_MRT4530\_62948C.1.pap  
US-10-437-963-164005

Query Match 34.0%; Score 33; DB 16; Length 39;  
Best Local Similarity 41.2%; Pred. No. 3.6e+02;  
Matches 7; Conservative 3; Mismatches 7; Indels 0; Gaps 0;

Qy 2 PNHLNFKIAFKIVSQEP 18  
Db 22 PPPLKNPAPFSLSHDP 38

RESULT 12  
US-10-424-599-208257  
; Sequence 208257, Application US/10424599  
; Publication No. US20040031072A1  
; GENERAL INFORMATION:  
; APPLICANT: La Rosa Thomas J  
; APPLICANT: Kovalic David K  
; APPLICANT: Zhou Yihua  
; APPLICANT: Cao Yongwei  
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With  
; FILE REFERENCE: 38-21(53223)B  
; CURRENT APPLICATION NUMBER: US/10/424,599  
; CURRENT FILING DATE: 2003-04-28  
; NUMBER OF SEQ ID NOS: 285684  
; SEQ ID NO 208257  
; LENGTH: 39  
; TYPE: PRT  
; ORGANISM: Glycine max  
; FEATURE:  
; OTHER INFORMATION: Clone ID: PAT\_MRT3847\_30081C.1.pap  
US-10-424-599-208257

Query Match 34.0%; Score 33; DB 15; Length 39;  
Best Local Similarity 50.0%; Pred. No. 3.7e+02;  
Matches 7; Conservative 1; Mismatches 6; Indels 0; Gaps 0;

Qy 4 HLNSKIAPKIVSQE 17  
 Db 26 HFTQSCFLIVSQE 39

## RESULT 13

US-10-424-599-199633  
 ; Sequence 199633, Application US/10424599  
 ; Publication No. US20040031072A1  
 ; GENERAL INFORMATION:  
 ; APPLICANT: La Rosa Thomas J  
 ; APPLICANT: Kovalic David K  
 ; APPLICANT: Zhou Yihua  
 ; APPLICANT: Cao Yongwei  
 ; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With  
 ; TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement  
 ; FILE REFERENCE: 38-21(53223)B  
 ; CURRENT APPLICATION NUMBER: US/10/424,599  
 ; CURRENT FILING DATE: 2003-04-28  
 ; NUMBER OF SEQ ID NOS: 285684  
 ; SEQ ID NO 199633  
 ; LENGTH: 46  
 ; TYPE: PRT  
 ; ORGANISM: Glycine max  
 ; FEATURE:  
 ; OTHER INFORMATION: Clone ID: PAT\_MRT3847\_22293C.1.pap  
 US-10-424-599-199633

Query Match 34.0%; Score 33; DB 15; Length 46;  
 Best Local Similarity 33.3%; Pred. No. 4.4e+02;  
 Matches 5; Conservative 4; Mismatches 6; Indels 0; Gaps 0;

Qy 3 NHLNSKIAPKIVSQE 17  
 Db 30 SHRNKRFLESIITSE 44

## RESULT 14

US-10-424-599-187478  
 ; Sequence 187478, Application US/10424599  
 ; Publication No. US20040031072A1  
 ; GENERAL INFORMATION:  
 ; APPLICANT: La Rosa Thomas J  
 ; APPLICANT: Kovalic David K  
 ; APPLICANT: Zhou Yihua  
 ; APPLICANT: Cao Yongwei  
 ; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With  
 ; TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement  
 ; FILE REFERENCE: 38-21(53223)B  
 ; CURRENT APPLICATION NUMBER: US/10/424,599  
 ; CURRENT FILING DATE: 2003-04-28  
 ; NUMBER OF SEQ ID NOS: 285684  
 ; SEQ ID NO 187478  
 ; LENGTH: 49  
 ; TYPE: PRT  
 ; ORGANISM: Glycine max  
 ; FEATURE:  
 ; OTHER INFORMATION: Clone ID: PAT\_MRT3847\_140304C.1.pap  
 US-10-424-599-187478

Query Match 34.0%; Score 33; DB 15; Length 49;  
 Best Local Similarity 41.7%; Pred. No. 4.7e+02;  
 Matches 5; Conservative 3; Mismatches 4; Indels 0; Gaps 0;

Qy 1 EPNHLSKIAPK 12  
 Db 8 EPKHNGRPSLK 19

## RESULT 15

US-09-864-761-35774  
 ; Sequence 35774, Application US/09864761  
 ; Patent No. US20020048763A1

; GENERAL INFORMATION:  
 ; APPLICANT: Penn, Sharon G.  
 ; APPLICANT: Rank, David R.  
 ; APPLICANT: Hanzel, David K.  
 ; APPLICANT: Chen, Wensheng  
 ; TITLE OF INVENTION: HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES USEFUL FOR  
 ; TITLE OF INVENTION: GENE EXPRESSION ANALYSIS BY MICROARRAY  
 ; FILE REFERENCE: Aemica-X-1  
 ; CURRENT APPLICATION NUMBER: US/09/864,761  
 ; CURRENT FILING DATE: 2001-05-23  
 ; PRIOR APPLICATION NUMBER: US 60/180,312  
 ; PRIOR FILING DATE: 2000-02-04  
 ; PRIOR APPLICATION NUMBER: US 60/207,456  
 ; PRIOR FILING DATE: 2000-05-26  
 ; PRIOR APPLICATION NUMBER: US 09/632,366  
 ; PRIOR FILING DATE: 2000-08-03  
 ; PRIOR APPLICATION NUMBER: GB 24263.6  
 ; PRIOR FILING DATE: 2000-10-04  
 ; PRIOR APPLICATION NUMBER: US 60/236,359  
 ; PRIOR FILING DATE: 2000-09-27  
 ; PRIOR APPLICATION NUMBER: PCT/US01/00666  
 ; PRIOR FILING DATE: 2001-01-30  
 ; PRIOR APPLICATION NUMBER: PCT/US01/00667  
 ; PRIOR FILING DATE: 2001-01-30  
 ; PRIOR APPLICATION NUMBER: PCT/US01/00664  
 ; PRIOR FILING DATE: 2001-01-30  
 ; PRIOR APPLICATION NUMBER: PCT/US01/00659  
 ; PRIOR FILING DATE: 2001-01-30  
 ; PRIOR APPLICATION NUMBER: PCT/US01/00665  
 ; PRIOR FILING DATE: 2001-01-30  
 ; PRIOR APPLICATION NUMBER: PCT/US01/00668  
 ; PRIOR FILING DATE: 2001-01-30  
 ; PRIOR APPLICATION NUMBER: PCT/US01/00663  
 ; PRIOR FILING DATE: 2001-01-30  
 ; PRIOR APPLICATION NUMBER: PCT/US01/00662  
 ; PRIOR FILING DATE: 2001-01-30  
 ; PRIOR APPLICATION NUMBER: PCT/US01/00661  
 ; PRIOR FILING DATE: 2001-01-30  
 ; PRIOR APPLICATION NUMBER: PCT/US01/00670  
 ; PRIOR FILING DATE: 2001-01-30  
 ; PRIOR APPLICATION NUMBER: US 60/234,687  
 ; PRIOR FILING DATE: 2000-09-21  
 ; PRIOR APPLICATION NUMBER: US 09/608,408  
 ; PRIOR FILING DATE: 2000-06-30  
 ; PRIOR APPLICATION NUMBER: US 09/774,203  
 ; PRIOR FILING DATE: 2001-01-29  
 ; NUMBER OF SEQ ID NOS: 49117  
 ; SOFTWARE: Annomax Sequence Listing Engine vers. 1.1  
 ; SEQ ID NO 35774  
 ; LENGTH: 36  
 ; TYPE: PRT  
 ; ORGANISM: Homo sapiens  
 ; FEATURE:  
 ; OTHER INFORMATION: MAP TO AP000470.1  
 ; OTHER INFORMATION: EXPRESSED IN HBL100, SIGNAL = 1.4  
 ; OTHER INFORMATION: EXPRESSED IN HEART, SIGNAL = 1.2  
 ; OTHER INFORMATION: EXPRESSED IN LUNG, SIGNAL = 5.3  
 ; OTHER INFORMATION: EXPRESSED IN BONE MARROW, SIGNAL = 2.2  
 ; OTHER INFORMATION: EXPRESSED IN BRAIN, SIGNAL = 1.2  
 ; OTHER INFORMATION: EXPRESSED IN ADULT LIVER, SIGNAL = 3.9  
 ; OTHER INFORMATION: EXPRESSED IN FETAL LIVER, SIGNAL = 1  
 ; OTHER INFORMATION: EXPRESSED IN PLACENTA, SIGNAL = 1.6  
 ; OTHER INFORMATION: EXPRESSED IN HELA, SIGNAL = 2.4  
 ; OTHER INFORMATION: EST HUMAN HIT: AA722296.1, EVALUE 4.00e-03  
 ; OTHER INFORMATION: SWISSPROT HIT: P55859, EVALUE 6.20e+00  
 US-09-864-761-35774

Query Match 33.5%; Score 32.5; DB 9; Length 36;  
 Best Local Similarity 53.8%; Pred. No. 4.1e+02;  
 Matches 7; Conservative 2; Mismatches 3; Indels 1; Gaps 1;

Qy 1 EPNHLSKIAPK 13  
 ||||| : : : : :  
 ||||| : : : : :

Db 19 EPNH-NSLLVFFL 30

## RESULT 16

US-10-424-599-267189  
; Sequence 267189, Application US/10424599  
; Publication No. US20040031072A1

## GENERAL INFORMATION:

APPLICANT: La Rosa Thomas J  
APPLICANT: Kovalic David K  
APPLICANT: Zhou Yihua  
APPLICANT: Cao Yongwei

TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With  
TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement

FILE REFERENCE: 38-21(53223)B

CURRENT APPLICATION NUMBER: US/10/424,599

CURRENT FILING DATE: 2003-04-28

NUMBER OF SEQ ID NOS: 285684

SEQ ID NO 267189

LENGTH: 27

TYPE: PRT

ORGANISM: Glycine max

FEATURE:

OTHER INFORMATION: Clone ID: PAT\_MRT3847\_83292C.1.pap  
US-10-424-599-267189

## Query Match

Best Local Similarity 33.0%; Score 32; DB 15; Length 27;

Matches 6; Conservative 3; Mismatches 3; Indels 0; Gaps 0;

Qy 5 LNSKIAFKIVSQ 16

||| |::|::|

Db 8 LNFKVDYKLYSQ 19

## RESULT 17

US-10-425-114-68647  
; Sequence 68647, Application US/10425114  
; Publication No. US20040034888A1

## GENERAL INFORMATION:

APPLICANT: Liu, Jingdong

APPLICANT: Zhou, Yihua

APPLICANT: Kovalic, David K.

APPLICANT: Screen, Steven E

APPLICANT: Tabaska, Jack E

APPLICANT: Cao, Yongwei

TITLE OF INVENTION: Nucleic Acid Molecules and Other Molecules Associated With  
TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement

FILE REFERENCE: 38-21(53313)B

CURRENT APPLICATION NUMBER: US/10/425,114

CURRENT FILING DATE: 2003-04-28

NUMBER OF SEQ ID NOS: 73128

SEQ ID NO 68647

LENGTH: 43

TYPE: PRT

ORGANISM: Zea mays

FEATURE:

OTHER INFORMATION: Clone ID: UC-ZMFLM017310A12\_FLI.pap  
US-10-425-114-68647

## Query Match

Best Local Similarity 33.0%; Score 32; DB 15; Length 43;

Matches 7; Conservative 3; Mismatches 8; Indels 0; Gaps 0;

Qy 1 EPNHLSKIAFKIVSQEP 18

||| |::|::|

Db 7 QPPSLAKIRHRLSKQTP 24

## RESULT 18

US-10-437-963-131784

; Sequence 131784, Application US/10437963

; Publication No. US20040123343A1

## GENERAL INFORMATION:

APPLICANT: La Rosa, Thomas J.

APPLICANT: Kovalic, David K.

APPLICANT: Zhou, Yihua

APPLICANT: Cao, Yongwei

APPLICANT: Wu, Wei

APPLICANT: Boukharov, Andrey A.

APPLICANT: Barbazuk, Brad

APPLICANT: Li, Ping

TITLE OF INVENTION: Rice Nucleic Acid Molecules and Other Molecules Associated With  
TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement

FILE REFERENCE: 38-21(53221)B

CURRENT APPLICATION NUMBER: US/10/437,963

CURRENT FILING DATE: 2003-05-14

NUMBER OF SEQ ID NOS: 204966

SEQ ID NO 131784

LENGTH: 43

TYPE: PRT

ORGANISM: Oryza sativa

FEATURE:

OTHER INFORMATION: Clone ID: PAT\_MRT4530\_33817C.1.pap  
US-10-437-963-131784

## Query Match

Best Local Similarity 33.0%; Score 32; DB 16; Length 43;

Matches 6; Conservative 3; Mismatches 3; Indels 0; Gaps 0;

Qy 3 NHLNSKIAFKIV 14

||| |::|::|

Db 25 SHLQWSISFKIV 36

## RESULT 19

US-10-424-599-245137

; Sequence 245137, Application US/10424599

; Publication No. US20040031072A1

## GENERAL INFORMATION:

APPLICANT: La Rosa Thomas J

APPLICANT: Kovalic David K

APPLICANT: Zhou Yihua

APPLICANT: Cao Yongwei

TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With  
TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement

FILE REFERENCE: 38-21(53223)B

CURRENT APPLICATION NUMBER: US/10/424,599

CURRENT FILING DATE: 2003-04-28

NUMBER OF SEQ ID NOS: 285684

SEQ ID NO 245137

LENGTH: 45

TYPE: PRT

ORGANISM: Glycine max

FEATURE:

OTHER INFORMATION: Clone ID: PAT\_MRT3847\_6338C.1.pap  
US-10-424-599-245137

## Query Match

Best Local Similarity 33.0%; Score 32; DB 15; Length 45;

Matches 4; Conservative 3; Mismatches 2; Indels 0; Gaps 0;

Qy 1 EPNHLSKI 9

||| |::|::|

Db 2 QPNHINVSIV 10

## RESULT 20

US-10-424-599-250836

; Sequence 250836, Application US/10424599

; Publication No. US20040031072A1

## GENERAL INFORMATION:

APPLICANT: La Rosa Thomas J

APPLICANT: Kovalic David K

APPLICANT: Zhou Yihua

APPLICANT: Cao Yongwei

```
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With
; TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement
; FILE REFERENCE: 38-21(53223)B
; CURRENT APPLICATION NUMBER: US/10/424,599
; CURRENT FILING DATE: 2003-04-28
; NUMBER OF SEQ ID NOS: 285684
; SEQ ID NO 250836
; LENGTH: 47
; TYPE: PRT
; ORGANISM: Glycine max
; FEATURE:
; OTHER INFORMATION: EXPRESSED IN PLACENTA, SIGNAL = 4
; OTHER INFORMATION: EXPRESSED IN BONE MARROW, SIGNAL = 4.6
; OTHER INFORMATION: EXPRESSED IN HBL100, SIGNAL = 5.1
; OTHER INFORMATION: EXPRESSED IN FETAL LIVER, SIGNAL = 3.5
; OTHER INFORMATION: EXPRESSED IN ADULT LIVER, SIGNAL = 4.1
; OTHER INFORMATION: EXPRESSED IN BRAIN, SIGNAL = 8.5
; OTHER INFORMATION: EXPRESSED IN BT474, SIGNAL = 3.2
; OTHER INFORMATION: EXPRESSED IN LUNG, SIGNAL = 4.7
; OTHER INFORMATION: EXPRESSED IN HELA, SIGNAL = 6
; OTHER INFORMATION: EXPRESSED IN HEART, SIGNAL = 4.3
; OTHER INFORMATION: EST HUMAN HIT: BF109229.1, EVALUE 1.00e-17
; OTHER INFORMATION: EST HUMAN HIT: AI738554.1, EVALUE 6.00e-13
; OTHER INFORMATION: SWISSPROT HIT: P48551, EVALUE 2.00e-21
US-10-424-599-250836

Query Match 33.0%; Score 32; DB 15; Length 47;
Best Local Similarity 66.7%; Pred. No. 6.6e+02;
Matches 6; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

QY 11 FKIVSQEPA 19
|||:::|
Db 11 FKIVNETPA 19

RESULT 21
US-09-864-761-33571
; Sequence 33571, Application US/09864761
; Patent No. US20020048763A1
; GENERAL INFORMATION:
; APPLICANT: Penn, Sharon G.
; APPLICANT: Rank, David R.
; APPLICANT: Hanzel, David K.
; APPLICANT: Chen, Wensheng
; TITLE OF INVENTION: HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES USEFUL FOR
; TITLE OF INVENTION: GENE EXPRESSION ANALYSIS BY MICROARRAY
; FILE REFERENCE: Aeomica-X-1
; CURRENT APPLICATION NUMBER: US/09/864,761
; CURRENT FILING DATE: 2001-05-23
; PRIOR APPLICATION NUMBER: US 60/180,312
; PRIOR FILING DATE: 2000-02-04
; PRIOR APPLICATION NUMBER: US 60/207,456
; PRIOR FILING DATE: 2000-05-26
; PRIOR APPLICATION NUMBER: US 09/632,366
; PRIOR FILING DATE: 2000-08-03
; PRIOR APPLICATION NUMBER: GB 24263.6
; PRIOR FILING DATE: 2000-10-04
; PRIOR APPLICATION NUMBER: US 60/236,359
; PRIOR FILING DATE: 2000-09-27
; PRIOR APPLICATION NUMBER: PCT/US01/00666
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00667
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00664
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00669
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00665
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00668
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00663
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00662
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00661
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00670
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: US 60/234,687
; PRIOR FILING DATE: 2000-09-21
; PRIOR APPLICATION NUMBER: US 09/608,408
; PRIOR FILING DATE: 2000-06-30
; PRIOR APPLICATION NUMBER: US 09/774,203
; PRIOR FILING DATE: 2001-01-29
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; NUMBER OF SEQ ID NOS: 49117
; SOFTWARE: Anomax Sequence Listing Engine vers. 1.1
; SEQ ID NO 33571
; LENGTH: 48
; TYPE: PRT
; ORGANISM: Homo sapiens
; FEATURE:
; OTHER INFORMATION: MAP TO AP000111.1
; OTHER INFORMATION: EXPRESSED IN PLACENTA, SIGNAL = 4
; OTHER INFORMATION: EXPRESSED IN BONE MARROW, SIGNAL = 4.6
; OTHER INFORMATION: EXPRESSED IN HBL100, SIGNAL = 5.1
; OTHER INFORMATION: EXPRESSED IN FETAL LIVER, SIGNAL = 3.5
; OTHER INFORMATION: EXPRESSED IN ADULT LIVER, SIGNAL = 4.1
; OTHER INFORMATION: EXPRESSED IN BRAIN, SIGNAL = 8.5
; OTHER INFORMATION: EXPRESSED IN BT474, SIGNAL = 3.2
; OTHER INFORMATION: EXPRESSED IN LUNG, SIGNAL = 4.7
; OTHER INFORMATION: EXPRESSED IN HELA, SIGNAL = 6
; OTHER INFORMATION: EXPRESSED IN HEART, SIGNAL = 4.3
; OTHER INFORMATION: EST HUMAN HIT: BF109229.1, EVALUE 1.00e-17
; OTHER INFORMATION: EST HUMAN HIT: AI738554.1, EVALUE 6.00e-13
; OTHER INFORMATION: SWISSPROT HIT: P48551, EVALUE 2.00e-21
US-09-864-761-33571

Query Match 33.0%; Score 32; DB 9; Length 48;
Best Local Similarity 33.3%; Pred. No. 6.8e+02;
Matches 5; Conservative 4; Mismatches 6; Indels 0; Gaps 0;

QY 3 NHLNSKIAPKIVSQE 17
|||:::|
Db 14 NHINVMVKPFSIVEE 28

RESULT 22
US-10-424-599-233753
; Sequence 233753, Application US/10424599
; Publication No. US20040031072A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa Thomas J
; APPLICANT: Kovalic David K
; APPLICANT: Zhou Yihua
; APPLICANT: Cao Yongwei
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With
; TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement
; FILE REFERENCE: 38-21(53223)B
; CURRENT APPLICATION NUMBER: US/10/424,599
; CURRENT FILING DATE: 2003-04-28
; NUMBER OF SEQ ID NOS: 285684
; SEQ ID NO 233753
; LENGTH: 47
; TYPE: PRT
; ORGANISM: Glycine max
; FEATURE:
; OTHER INFORMATION: Clone ID: PAT_MRT3847_53103C.1.pap
US-10-424-599-233753

Query Match 32.5%; Score 31.5; DB 15; Length 47;
Best Local Similarity 33.3%; Pred. No. 8.1e+02;
Matches 6; Conservative 7; Mismatches 2; Indels 3; Gaps 1;

QY 2 PNLNSKIAP---KIVSQ 16
|||:::|
Db 9 PHIHSSLIYFPTKLVSQ 26

RESULT 23
US-10-084-813-333
; Sequence 333, Application US/10084813
; Publication No. US20030068615A1
; GENERAL INFORMATION:
; APPLICANT: SAXINGER, CARL
; TITLE OF INVENTION: POLYPEPTIDES THAT BIND HIV GP120 AND RELATED NUCLEIC
; TITLE OF INVENTION: ACIDS, ANTIBODIES, COMPOSITIONS, AND METHODS OF USE
; FILE REFERENCE: 2158/75
```

```
; CURRENT APPLICATION NUMBER: US/10/084,813
; CURRENT FILING DATE: 2002-02-27
; PRIOR APPLICATION NUMBER: PCT/US00/23505
; PRIOR FILING DATE: 2000-08-25
; PRIOR APPLICATION NUMBER: US 60/151,270
; PRIOR FILING DATE: 1999-08-27
; NUMBER OF SEQ ID NOS: 1242
; SOFTWARE: PatentIn version 3.1
; SEQ ID NO 333
; LENGTH: 21
; TYPE: PRT
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence: binding peptide
US-10-084-813-333

Query Match      32.0%; Score 31; DB 14; Length 21;
Best Local Similarity 38.5%; Pred. No. 4e+02;
Matches 5; Conservative 5; Mismatches 3; Indels 0; Gaps 0;

Qy      6 NSKIAFKIVSQEP 18
Db      8 NKEVSKRVITQDP 20

RESULT 24
US-10-084-813-334
; Sequence 334, Application US/10084813
; Publication No. US20030068615A1
; GENERAL INFORMATION:
; APPLICANT: SAXINGER, CARL
; TITLE OF INVENTION: POLYPEPTIDES THAT BIND HIV GP120 AND RELATED NUCLEIC
; FILE REFERENCE: 215875
; CURRENT APPLICATION NUMBER: US/10/084,813
; CURRENT FILING DATE: 2002-02-27
; PRIOR APPLICATION NUMBER: PCT/US00/23505
; PRIOR FILING DATE: 2000-08-25
; PRIOR APPLICATION NUMBER: US 60/151,270
; PRIOR FILING DATE: 1999-08-27
; NUMBER OF SEQ ID NOS: 1242
; SOFTWARE: PatentIn version 3.1
; SEQ ID NO 334
; LENGTH: 21
; TYPE: PRT
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence: binding peptide
US-10-084-813-334

Query Match      32.0%; Score 31; DB 14; Length 21;
Best Local Similarity 38.5%; Pred. No. 4e+02;
Matches 5; Conservative 5; Mismatches 3; Indels 0; Gaps 0;

Qy      6 NSKIAFKIVSQEP 18
Db      3 NKEVSKRVITQDP 15

RESULT 25
US-10-437-963-150845
; Sequence 150845, Application US/10437963
; Publication No. US20040123343A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa, Thomas J.
; APPLICANT: Kovalic, David K.
; APPLICANT: Zhou, Yihua
; APPLICANT: Cao, Yongwei
; APPLICANT: Wu, Wei
; APPLICANT: Boukharov, Andrey A.
; APPLICANT: Barbazov, Brad
; APPLICANT: Li, Ping
; TITLE OF INVENTION: Rice Nucleic Acid Molecules and Other Molecules Associated With

; CURRENT APPLICATION NUMBER: US/10/084,813
; CURRENT FILING DATE: 2002-02-27
; PRIOR APPLICATION NUMBER: PCT/US00/23505
; PRIOR FILING DATE: 2000-08-25
; PRIOR APPLICATION NUMBER: US 60/151,270
; PRIOR FILING DATE: 1999-08-27
; NUMBER OF SEQ ID NOS: 1242
; SOFTWARE: PatentIn version 3.1
; SEQ ID NO 333
; LENGTH: 21
; TYPE: PRT
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence: binding peptide
US-10-084-813-333

Query Match      32.0%; Score 31; DB 14; Length 21;
Best Local Similarity 38.5%; Pred. No. 4e+02;
Matches 5; Conservative 5; Mismatches 3; Indels 0; Gaps 0;

Qy      6 NSKIAFKIVSQEP 18
Db      3 NKEVSKRVITQDP 15

RESULT 25
US-10-437-963-150845
; Sequence 150845, Application US/10437963
; Publication No. US20040123343A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa, Thomas J.
; APPLICANT: Kovalic, David K.
; APPLICANT: Zhou, Yihua
; APPLICANT: Cao, Yongwei
; APPLICANT: Wu, Wei
; APPLICANT: Boukharov, Andrey A.
; APPLICANT: Barbazov, Brad
; APPLICANT: Li, Ping
; TITLE OF INVENTION: Rice Nucleic Acid Molecules and Other Molecules Associated With

; TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement
; FILE REFERENCE: 38-21(53221)B
; CURRENT APPLICATION NUMBER: US/10/437,963
; CURRENT FILING DATE: 2003-05-14
; NUMBER OF SEQ ID NOS: 204966
; SEQ ID NO 150845
; LENGTH: 30
; TYPE: PRT
; ORGANISM: Oryza sativa
; FEATURE:
; OTHER INFORMATION: Clone ID: PAT_MRT4530_51044C.1.pep
US-10-437-963-150845

Query Match      32.0%; Score 31; DB 16; Length 30;
Best Local Similarity 50.0%; Pred. No. 5.9e+02;
Matches 7; Conservative 1; Mismatches 6; Indels 0; Gaps 0;

Qy      6 NSKIAFKIVSQEPA 19
Db      4 NSHAYFKTVSNPPS 17

RESULT 26
US-09-801-348-29
; Sequence 29, Application US/09801348
; Publication No. US20040166530A1
; GENERAL INFORMATION:
; APPLICANT: Harper, Jeffrey W.
; APPLICANT: Elledge, Stephen J.
; TITLE OF INVENTION: F-BOX PROTEINS AND GENES
; FILE REFERENCE: BCM-03510
; CURRENT APPLICATION NUMBER: US/09/801,348
; CURRENT FILING DATE: 2001-07-31
; EARLIER APPLICATION NUMBER: 09/172,841
; EARLIER FILING DATE: 1998-10-15
; EARLIER APPLICATION NUMBER: 08/951,621
; EARLIER FILING DATE: 1997-10-16
; NUMBER OF SEQ ID NOS: 60
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 29
; LENGTH: 33
; TYPE: PRT
; ORGANISM: Mus musculus
; US-09-801-348-29

Query Match      32.0%; Score 31; DB 11; Length 33;
Best Local Similarity 58.3%; Pred. No. 6.6e+02;
Matches 7; Conservative 2; Mismatches 3; Indels 0; Gaps 0;

Qy      5 LNSKIAFKIVSQ 16
Db      1 LPABITFKIFSQ 12

RESULT 27
US-09-764-891-4356
; Sequence 4356, Application US/09764891
; Publication No. US20030077808A1
; GENERAL INFORMATION:
; APPLICANT: Rosen et al.
; TITLE OF INVENTION: Nucleic Acids, Proteins, and Antibodies
; FILE REFERENCE: PC006
; CURRENT APPLICATION NUMBER: US/09/764,891
; CURRENT FILING DATE: 2001-01-17
; Prior application data removed - consult PALM or file wrapper
; NUMBER OF SEQ ID NOS: 10231
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 4356
; LENGTH: 34
; TYPE: PRT
; ORGANISM: Homo sapiens
; FEATURE:
; NAME/KEY: SITE
```

```
; LOCATION: (3)
; OTHER INFORMATION: Xaa equals any of the naturally occurring L-amino acids
; NAME/KEY: SITE
; LOCATION: (20)
; OTHER INFORMATION: Xaa equals any of the naturally occurring L-amino acids
US-09-764-891-4356

Query Match          32.0%; Score 31; DB 10; Length 34;
Best Local Similarity 40.0%; Pred. No. 6.8e+02;
Matches 6; Conservative 3; Mismatches 6; Indels 0; Gaps 0;

QY      3 NHLNSKIAPKIVSQE 17
Db      12 HHLNQVIXNLIISNK 26

RESULT 28
US-10-062-831-72
; Sequence 72, Application US/10062831
; Publication No. US20030105297A1
; GENERAL INFORMATION:
; APPLICANT: Steven M. Ruben, et al.
; TITLE OF INVENTION: 32 Human Secreted Proteins
; FILE REFERENCE: PZ006P1
; CURRENT APPLICATION NUMBER: US/10/062,831
; CURRENT FILING DATE: 2002-02-05
; PRIOR APPLICATION NUMBER: 09/690,454
; PRIOR FILING DATE: 1998-11-10
; PRIOR APPLICATION NUMBER: PCT/US98/10869
; PRIOR FILING DATE: May 28, 1998
; PRIOR APPLICATION NUMBER: 60/044,039
; PRIOR FILING DATE: May 30, 1997
; PRIOR APPLICATION NUMBER: 60/048,093
; PRIOR FILING DATE: May 30, 1997
; PRIOR APPLICATION NUMBER: 60/048,190
; PRIOR FILING DATE: May 30, 1997
; PRIOR APPLICATION NUMBER: 60/050,935
; PRIOR FILING DATE: May 30, 1997
; PRIOR APPLICATION NUMBER: 60/048,101
; PRIOR FILING DATE: May 30, 1997
; PRIOR APPLICATION NUMBER: 60/048,356
; PRIOR FILING DATE: May 30, 1997
; PRIOR APPLICATION NUMBER: 60/056,250
; PRIOR FILING DATE: August 29, 1997
; PRIOR APPLICATION NUMBER: 60/056,296
; PRIOR FILING DATE: August 29, 1997
; PRIOR APPLICATION NUMBER: 60/056,293
; PRIOR FILING DATE: August 29, 1997
; NUMBER OF SEQ ID NOS: 229
; SOFTWARE: Patentin Ver. 2.0
; SEQ ID NO 72
; LENGTH: 34
; TYPE: PRT
; ORGANISM: Homo sapiens
; FEATURE:
; NAME/KEY: SITE
; LOCATION: (34)
; OTHER INFORMATION: Xaa equals stop translation
US-10-062-831-72

Query Match          32.0%; Score 31; DB 14; Length 34;
Best Local Similarity 47.4%; Pred. No. 6.8e+02;
Matches 9; Conservative 2; Mismatches 4; Indels 4; Gaps 1;

QY      5 LNSKIAPKIV-----SQEPA 19
Db      11 LNSKLVAAVVNLKASQMPA 29

RESULT 29
US-10-205-428-383
; Sequence 383, Application US/10205428
; Publication No. US20030108907A1
```

```
; GENERAL INFORMATION:
; APPLICANT: Rosen et al.
; TITLE OF INVENTION: Nucleic Acids, Proteins, and Antibodies
; FILE REFERENCE: PAIL7C1
; CURRENT APPLICATION NUMBER: US/10/205,428
; CURRENT FILING DATE: 2002-07-26
; PRIOR APPLICATION NUMBER: 09/764,892
; PRIOR FILING DATE: 2001-01-17
; PRIOR APPLICATION NUMBER: 60/179,065
; PRIOR FILING DATE: 2000-01-31
; PRIOR APPLICATION NUMBER: 60/180,628
; PRIOR FILING DATE: 2000-02-04
; PRIOR APPLICATION NUMBER: 60/214,886
; PRIOR FILING DATE: 2000-06-28
; PRIOR APPLICATION NUMBER: 60/217,487
; PRIOR FILING DATE: 2000-07-11
; PRIOR APPLICATION NUMBER: 60/225,758
; PRIOR FILING DATE: 2000-08-14
; PRIOR APPLICATION NUMBER: 60/220,963
; PRIOR FILING DATE: 2000-07-26
; PRIOR APPLICATION NUMBER: 60/217,496
; PRIOR FILING DATE: 2000-07-11
; PRIOR APPLICATION NUMBER: 60/225,447
; PRIOR FILING DATE: 2000-08-14
; PRIOR APPLICATION NUMBER: 60/218,290
; PRIOR FILING DATE: 2000-07-14
; Remaining Prior Application data removed - See File Wrapper or PALM.
; NUMBER OF SEQ ID NOS: 1019
; SOFTWARE: Patentin Ver. 2.0
; SEQ ID NO 383
; LENGTH: 34
; TYPE: PRT
; ORGANISM: Homo sapiens
; FEATURE:
; NAME/KEY: misc_feature
; LOCATION: (3)
; OTHER INFORMATION: Xaa equals any of the naturally occurring L-amino acids
; FEATURE:
; NAME/KEY: misc_feature
; LOCATION: (20)
; OTHER INFORMATION: Xaa equals any of the naturally occurring L-amino acids
US-10-205-428-383

Query Match          32.0%; Score 31; DB 14; Length 34;
Best Local Similarity 40.0%; Pred. No. 6.8e+02;
Matches 6; Conservative 3; Mismatches 6; Indels 0; Gaps 0;

QY      3 NHLNSKIAPKIVSQE 17
Db      12 HHLNQVIXNLIISNK 26

RESULT 30
US-10-062-599-72
; Sequence 72, Application US/10062599
; Publication No. US20030195346A1
; GENERAL INFORMATION:
; APPLICANT: Steven M. Ruben, et al.
; TITLE OF INVENTION: 32 Human Secreted Proteins
; FILE REFERENCE: PZ006P1
; CURRENT APPLICATION NUMBER: US/10/062,599
; CURRENT FILING DATE: 2002-02-05
; PRIOR APPLICATION NUMBER: 09/690,454
; PRIOR FILING DATE: 2000-10-18
; PRIOR APPLICATION NUMBER: 09/189,144
; PRIOR FILING DATE: 1998-11-10
; PRIOR APPLICATION NUMBER: 60/044,039
; PRIOR FILING DATE: May 30, 1997
; PRIOR APPLICATION NUMBER: 60/048,093
; PRIOR FILING DATE: May 30, 1997
; PRIOR APPLICATION NUMBER: 60/048,190
; PRIOR FILING DATE: May 30, 1997
; PRIOR APPLICATION NUMBER: 60/050,935
```

```
; PRIOR FILING DATE: May 30, 1997
; PRIOR APPLICATION NUMBER: 60/048,101
; PRIOR FILING DATE: May 30, 1997
; PRIOR APPLICATION NUMBER: 60/048,356
; PRIOR FILING DATE: May 30, 1997
; PRIOR APPLICATION NUMBER: 60/056,250
; PRIOR FILING DATE: August 29, 1997
; PRIOR APPLICATION NUMBER: 60/056,296
; PRIOR FILING DATE: August 29, 1997
; PRIOR APPLICATION NUMBER: 60/056,293
; PRIOR FILING DATE: August 29, 1997
; NUMBER OF SEQ ID NOS: 229
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 72
; LENGTH: 34
; TYPE: PRT
; ORGANISM: Homo sapiens
; FEATURE:
; NAME/KEY: SITE
; LOCATION: (34)
; OTHER INFORMATION: xaa equals stop translation
US-10-062-599-72
```

```
Query Match 32.0%; Score 31; DB 14; Length 34;
Best Local Similarity 47.4%; Pred. No. 6.8e+02;
Matches 9; Conservative 2; Mismatches 4; Indels 4; Gaps 1;
```

```
Qy 5 LNSKIAFKIV---SQEFA 19
| | | | : | | |
Db 11 LNSKLVAAVVNLKASQMPA 29
```

```
RESULT 31
US-10-424-599-227023
; Sequence 227023, Application US/10424599
; Publication No. US20040031072A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa Thomas J
; APPLICANT: Kovalic David K
; APPLICANT: Zhou Yihua
; APPLICANT: Cao Yongwei
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With
; FILE OF INVENTION: Plants and Uses Thereof for Plant Improvement
; FILE REFERENCE: 38-21(53223)B
; CURRENT APPLICATION NUMBER: US/10/424,599
; CURRENT FILING DATE: 2003-04-28
; NUMBER OF SEQ ID NOS: 285684
; SEQ ID NO 227023
; LENGTH: 42
; TYPE: PRT
; ORGANISM: Glycine max
; FEATURE:
; OTHER INFORMATION: Clone ID: PAT_MRT3847_47031C.1.pep
US-10-424-599-227023
```

```
Query Match 32.0%; Score 31; DB 15; Length 42;
Best Local Similarity 50.0%; Pred. No. 8.6e+02;
Matches 5; Conservative 3; Mismatches 2; Indels 0; Gaps 0;
```

```
Qy 3 NHLNSKIATFK 12
| | | : | | |
Db 26 NHHGDIATFE 35
```

```
RESULT 32
US-10-029-386-33676
; Sequence 33676, Application US/10029386
; Publication No. US20030194704A1
; GENERAL INFORMATION:
; APPLICANT: Penn, Sharron G.
; APPLICANT: Rank, David R.
; APPLICANT: Hanzel, David K.
; TITLE OF INVENTION: HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES USEFUL FOR G
```

```
; TITLE OF INVENTION: EXPRESSION ANALYSIS TWO
; FILE REFERENCE: AEOMICA-X-2
; CURRENT APPLICATION NUMBER: US/10/029,386
; CURRENT FILING DATE: 2001-12-20
; NUMBER OF SEQ ID NOS: 34288
; SOFTWARE: Annomax Sequence Listing Engine vers. 1.1
; SEQ ID NO 33676
; LENGTH: 43
; TYPE: PRT
; ORGANISM: Homo sapiens
; FEATURE:
; OTHER INFORMATION: MAP TO AC004030.1
; OTHER INFORMATION: EXPRESSED IN HELA, SIGNAL = 0.42
US-10-029-386-33676
```

```
Query Match 32.0%; Score 31; DB 14; Length 43;
Best Local Similarity 41.7%; Pred. No. 8.8e+02;
Matches 5; Conservative 4; Mismatches 3; Indels 0; Gaps 0;
```

```
Qy 2 PNHLSNKIAFKI 13
| | | : | | |
Db 15 PHLHSNVAVTV 26
```

```
RESULT 33
US-09-864-761-41924
; Sequence 41924, Application US/09864761
; Patent No. US20020048763A1
; GENERAL INFORMATION:
; APPLICANT: Penn, Sharron G.
; APPLICANT: Rank, David R.
; APPLICANT: Hanzel, David K.
; APPLICANT: Chen, Wensheng
; TITLE OF INVENTION: HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES USEFUL FOR
; FILE REFERENCE: Aeomica-X-1
; CURRENT APPLICATION NUMBER: US/09/864,761
; CURRENT FILING DATE: 2001-05-23
; PRIOR APPLICATION NUMBER: US 60/180,312
; PRIOR FILING DATE: 2000-02-04
; PRIOR APPLICATION NUMBER: US 60/207,456
; PRIOR FILING DATE: 2000-05-26
; PRIOR APPLICATION NUMBER: US 09/632,366
; PRIOR FILING DATE: 2000-08-03
; PRIOR APPLICATION NUMBER: GB 24263.6
; PRIOR FILING DATE: 2000-10-04
; PRIOR APPLICATION NUMBER: US 60/236,359
; PRIOR FILING DATE: 2000-09-27
; PRIOR APPLICATION NUMBER: PCT/US01/00666
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00667
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00664
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00669
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00665
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00668
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00663
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00662
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00661
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00670
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: US 60/234,687
; PRIOR FILING DATE: 2000-09-21
; PRIOR APPLICATION NUMBER: US 09/608,408
; PRIOR FILING DATE: 2000-06-30
; PRIOR APPLICATION NUMBER: US 09/774,203
```

```
; PRIOR FILING DATE: 2001-01-29
; NUMBER OF SEQ ID NOS: 49117
; SOFTWARE: Anomax Sequence Listing Engine vers. 1.1
; SEQ ID NO 41924
; LENGTH: 44
; TYPE: PRT
; ORGANISM: Homo sapiens
; FEATURE:
; OTHER INFORMATION: MAP TO AC022211.2
; OTHER INFORMATION: EXPRESSED IN ADULT LIVER, SIGNAL = 1.5
; OTHER INFORMATION: EXPRESSED IN HELA, SIGNAL = 2.6
; OTHER INFORMATION: EXPRESSED IN HEART, SIGNAL = 2.3
; OTHER INFORMATION: EXPRESSED IN BRAIN, SIGNAL = 3.5
; OTHER INFORMATION: EXPRESSED IN PLACENTA, SIGNAL = 2.8
; OTHER INFORMATION: EXPRESSED IN BONE MARROW, SIGNAL = 3.4
; OTHER INFORMATION: EXPRESSED IN LUNG, SIGNAL = 3.9
; OTHER INFORMATION: EXPRESSED IN FETAL LIVER, SIGNAL = 3.4
; OTHER INFORMATION: SWISSPROT HIT: P55855, EVALUE 3.00e-20
; OTHER INFORMATION: EST_HUMAN HIT: BF574192.1, EVALUE 4.00e-19
US-09-864-761-41924
```

```
Query Match 32.0%; Score 31; DB 9; Length 44;
Best Local Similarity 33.3%; Pred. No. 9.1e+02;
Matches 8; Conservative 3; Mismatches 5; Indels 8; Gaps 1;
```

```
QY 3 NHLNSKIA-----FKIVSQEP 18
Db 9 DHNLKVGQDGSVVQFKLRHTP 32
```

```
RESULT 34
US-10-424-599-181955
; Sequence 181955, Application US/10424599
; Publication No. US20040031072A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa Thomas J
; APPLICANT: Kovalic David K
; APPLICANT: Zhou Yihua
; APPLICANT: Cao Yongwei
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With
; FILE REFERENCE: 38-21(53223)B
; CURRENT APPLICATION NUMBER: US/10/424,599
; CURRENT FILING DATE: 2003-04-28
; NUMBER OF SEQ ID NOS: 285684
; SEQ ID NO 181955
; LENGTH: 46
; TYPE: PRT
; ORGANISM: Glycine max
; FEATURE:
; OTHER INFORMATION: Clone ID: PAT_MRT3847_135319C.1.pcp
US-10-424-599-181955
```

```
Query Match 32.0%; Score 31; DB 15; Length 46;
Best Local Similarity 44.4%; Pred. No. 9.5e+02;
Matches 8; Conservative 2; Mismatches 8; Indels 0; Gaps 0;
```

```
QY 2 PNHLSKIAFKIVSQEPA 19
Db 27 PIHVNNGCATKIESSPA 44
```

```
RESULT 35
US-09-984-429-87
; Sequence 87, Application US/09984429
; Publication No. US20040010132A1
; GENERAL INFORMATION:
; APPLICANT: Rosen et al.
; TITLE OF INVENTION: 53 Human Secreted Proteins
; FILE REFERENCE: P2018P2
; CURRENT APPLICATION NUMBER: US/09/984,429
; CURRENT FILING DATE: 2001-10-30
; PRIOR APPLICATION NUMBER: 60/244,591
```

```
; PRIOR FILING DATE: 2000-11-01
; PRIOR APPLICATION NUMBER: 09/288,143
; PRIOR FILING DATE: 1999-04-08
; PRIOR APPLICATION NUMBER: PCT/US98/21142
; PRIOR FILING DATE: 1998-10-08
; PRIOR APPLICATION NUMBER: 60/061,463
; PRIOR FILING DATE: 1997-10-09
; PRIOR APPLICATION NUMBER: 60/061,529
; PRIOR FILING DATE: 1997-10-09
; PRIOR APPLICATION NUMBER: 60/071,498
; PRIOR FILING DATE: 1997-10-09
; PRIOR APPLICATION NUMBER: 60/061,527
; PRIOR FILING DATE: 1997-10-09
; PRIOR APPLICATION NUMBER: 60/061,536
; PRIOR FILING DATE: 1997-10-09
; PRIOR APPLICATION NUMBER: 60/061,532
; PRIOR FILING DATE: 1997-10-09
; NUMBER OF SEQ ID NOS: 727
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 87
; LENGTH: 47
; TYPE: PRT
; ORGANISM: Homo sapiens
US-09-984-429-87
```

```
Query Match 32.0%; Score 31; DB 11; Length 47;
Best Local Similarity 45.5%; Pred. No. 9.8e+02;
Matches 5; Conservative 4; Mismatches 2; Indels 0; Gaps 0;
```

```
QY 9 IAFKIVSQEPA 19
Db 19 VAFRLTNQIPA 29
```

```
RESULT 36
US-10-393-840-84
; Sequence 84, Application US/10393840
; Publication No. US20030229922A1
; GENERAL INFORMATION:
; APPLICANT: Bloksberg, Leonard N.
; TITLE OF INVENTION: Materials and Methods for the
; FILE REFERENCE: 11000.1012c3
; CURRENT APPLICATION NUMBER: US/10/393,840
; CURRENT FILING DATE: 2003-03-20
; PRIOR APPLICATION NUMBER: US 09/636,800
; PRIOR FILING DATE: 2000-08-10
; PRIOR APPLICATION NUMBER: US 09/170,862
; PRIOR FILING DATE: 1998-10-13
; PRIOR APPLICATION NUMBER: US 60/148,426
; PRIOR FILING DATE: 1999-08-11
; PRIOR APPLICATION NUMBER: PCT NZ/99/00169
; PRIOR FILING DATE: 1999-10-08
; NUMBER OF SEQ ID NOS: 956
; SOFTWARE: FastSeq for Windows Version 4.0
; SEQ ID NO 84
; LENGTH: 47
; TYPE: PRT
; ORGANISM: Eucalyptus grandis
US-10-393-840-84
```

```
Query Match 32.0%; Score 31; DB 14; Length 47;
Best Local Similarity 50.0%; Pred. No. 9.8e+02;
Matches 7; Conservative 3; Mismatches 4; Indels 0; Gaps 0;
```

```
QY 5 INSKIAFKIVSQEP 18
Db 4 LDSADAFKSVRRDP 17
```

```
RESULT 37
US-10-424-599-223920
; Sequence 223920, Application US/10424599
```



```
; Publication No. US20040031072A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa Thomas J
; APPLICANT: Kovalic David K
; APPLICANT: Zhou Yihua
; APPLICANT: Cao Yongwei
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With
; FILE OF INVENTION: Plants and Uses Thereof for Plant Improvement
; FILE REFERENCE: 38-21(53223)B
; CURRENT APPLICATION NUMBER: US/10/424,599
; CURRENT FILING DATE: 2003-04-28
; NUMBER OF SEQ ID NOS: 285684
; SEQ ID NO 223920
; LENGTH: 47
; TYPE: PRT
; ORGANISM: Glycine max
; FEATURE:
; OTHER INFORMATION: Clone ID: PAT_MRT3847_44229C.1.pep
US-10-424-599-223920

Query Match          32.0%; Score 31; DB 15; Length 47;
Best Local Similarity 31.2%; Pred. No. 9.8e+02;
Matches 5; Conservative 4; Mismatches 7; Indels 0; Gaps 0;

QY 3 NHLNSKIAFKIVSQEP 18
|::||::|
Db 19 NFLNRAEAVDCIGEP 34

RESULT 38
US-10-150-111-87
; Sequence 87, Application US/10150111
; Publication No. US20030078386A1
; GENERAL INFORMATION:
; APPLICANT: Rubin et al.
; TITLE OF INVENTION: Secreted Protein HFEAD48
; FILE REFERENCE: PZ018PDI
; CURRENT APPLICATION NUMBER: US/10/150,111
; CURRENT FILING DATE: 2002-05-20
; PRIOR APPLICATION NUMBER: 09/288,143
; PRIOR FILING DATE: 1999-04-08
; PRIOR APPLICATION NUMBER: PCT/US98/21142
; PRIOR FILING DATE: 1998-10-08
; PRIOR APPLICATION NUMBER: 60/061,463
; PRIOR FILING DATE: 1997-10-09
; PRIOR APPLICATION NUMBER: 60/061,529
; PRIOR FILING DATE: 1997-10-09
; PRIOR APPLICATION NUMBER: 60/071,498
; PRIOR FILING DATE: 1997-10-09
; PRIOR APPLICATION NUMBER: 60/061,527
; PRIOR FILING DATE: 1997-10-09
; PRIOR APPLICATION NUMBER: 60/061,536
; PRIOR FILING DATE: 1997-10-09
; PRIOR APPLICATION NUMBER: 60/061,532
; PRIOR FILING DATE: 1997-10-09
; NUMBER OF SEQ ID NOS: 219
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 87
; LENGTH: 48
; TYPE: PRT
; ORGANISM: Homo sapiens
; FEATURE:
; NAME/KEY: SITE
; LOCATION: (48)
; OTHER INFORMATION: Xaa equals stop translation
US-10-150-111-87

Query Match          32.0%; Score 31; DB 14; Length 48;
Best Local Similarity 45.5%; Pred. No. 1e+03;
Matches 5; Conservative 4; Mismatches 2; Indels 0; Gaps 0;

QY 9 IAFKIVSQEPA 19
|::||::|

; Publication No. US20040031072A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa Thomas J
; APPLICANT: Kovalic David K
; APPLICANT: Zhou Yihua
; APPLICANT: Cao Yongwei
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With
; FILE OF INVENTION: Plants and Uses Thereof for Plant Improvement
; FILE REFERENCE: 38-21(53223)B
; CURRENT APPLICATION NUMBER: US/10/424,599
; CURRENT FILING DATE: 2003-04-28
; NUMBER OF SEQ ID NOS: 285684
; SEQ ID NO 223920
; LENGTH: 47
; TYPE: PRT
; ORGANISM: Glycine max
; FEATURE:
; OTHER INFORMATION: Clone ID: PAT_MRT3847_44229C.1.pep
US-10-424-599-223920

Db 19 VAFRLTNQIPA 29

RESULT 39
US-10-424-599-156398
; Sequence 156398, Application US/10424599
; Publication No. US20040031072A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa Thomas J
; APPLICANT: Kovalic David K
; APPLICANT: Zhou Yihua
; APPLICANT: Cao Yongwei
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With
; FILE OF INVENTION: Plants and Uses Thereof for Plant Improvement
; FILE REFERENCE: 38-21(53223)B
; CURRENT APPLICATION NUMBER: US/10/424,599
; CURRENT FILING DATE: 2003-04-28
; NUMBER OF SEQ ID NOS: 285684
; SEQ ID NO 156398
; LENGTH: 48
; TYPE: PRT
; ORGANISM: Glycine max
; FEATURE:
; OTHER INFORMATION: Clone ID: PAT_MRT3847_112248C.1.pep
US-10-424-599-156398

Query Match          32.0%; Score 31; DB 15; Length 48;
Best Local Similarity 37.5%; Pred. No. 1e+03;
Matches 6; Conservative 4; Mismatches 6; Indels 0; Gaps 0;

QY 3 NHLNSKIAFKIVSQEP 18
|::||::|
Db 9 NNIYSKLLFSLSSUPP 24

RESULT 40
US-09-764-860-359
; Sequence 359, Application US/09764860
; Patent No. US20020094953A1
; GENERAL INFORMATION:
; APPLICANT: Rosen et al.
; TITLE OF INVENTION: Nucleic Acids, Proteins, and Antibodies
; FILE REFERENCE: PC008
; CURRENT APPLICATION NUMBER: US/09/764,860
; CURRENT FILING DATE: 2001-01-17
; Prior application data removed - consult PALM or file wrapper
; NUMBER OF SEQ ID NOS: 1198
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 359
; LENGTH: 50
; TYPE: PRT
; ORGANISM: Homo sapiens
; OTHER INFORMATION:
US-09-764-860-359

Query Match          32.0%; Score 31; DB 9; Length 50;
Best Local Similarity 60.0%; Pred. No. 1e+03;
Matches 6; Conservative 1; Mismatches 3; Indels 0; Gaps 0;

QY 2 PNHLNSKIAF 11
|::||::|
Db 36 PPHVNWKTAF 45

RESULT 41
US-10-074-095-359
; Sequence 359, Application US/10074095
; Publication No. US20030077704A1
; GENERAL INFORMATION:
; APPLICANT: Rosen et al.
; TITLE OF INVENTION: Nucleic Acids, Proteins, and Antibodies
; FILE REFERENCE: PC008C1
; CURRENT APPLICATION NUMBER: US/10/074,095
; CURRENT FILING DATE: 2002-02-14
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1 ; PRIOR APPLICATION NUMBER: 09/764,860  
2 ; PRIOR FILING DATE: 2001-01-17  
3 ; PRIOR APPLICATION NUMBER: 60/179,065  
4 ; PRIOR FILING DATE: 2000-01-31  
5 ; PRIOR APPLICATION NUMBER: 60/180,628  
6 ; PRIOR FILING DATE: 2000-02-04  
7 ; PRIOR APPLICATION NUMBER: 60/214,886  
8 ; PRIOR FILING DATE: 2000-06-28  
9 ; PRIOR APPLICATION NUMBER: 60/217,487  
10 ; PRIOR FILING DATE: 2000-07-11  
11 ; PRIOR APPLICATION NUMBER: 60/225,758  
12 ; PRIOR FILING DATE: 2000-08-14  
13 ; PRIOR APPLICATION NUMBER: 60/220,963  
14 ; PRIOR FILING DATE: 2000-07-26  
15 ; PRIOR APPLICATION NUMBER: 60/217,496  
16 ; PRIOR FILING DATE: 2000-07-11  
17 ; PRIOR APPLICATION NUMBER: 60/225,447  
18 ; PRIOR FILING DATE: 2000-08-14  
19 ; PRIOR APPLICATION NUMBER: 60/218,290  
20 ; PRIOR FILING DATE: 2000-07-14  
21 ; PRIOR APPLICATION NUMBER: 60/225,757  
22 ; PRIOR FILING DATE: 2000-08-14  
23 ; PRIOR APPLICATION NUMBER: 60/226,868  
24 ; PRIOR FILING DATE: 2000-08-22  
25 ; PRIOR APPLICATION NUMBER: 60/216,647  
26 ; PRIOR FILING DATE: 2000-07-07  
27 ; PRIOR APPLICATION NUMBER: 60/225,267  
28 ; PRIOR FILING DATE: 2000-08-14  
29 ; PRIOR APPLICATION NUMBER: 60/216,880  
30 ; PRIOR FILING DATE: 2000-07-07  
31 ; PRIOR APPLICATION NUMBER: 60/225,270  
32 ; PRIOR FILING DATE: 2000-08-14  
33 ; PRIOR APPLICATION NUMBER: 60/251,869  
34 ; PRIOR FILING DATE: 2000-12-08  
35 ; PRIOR APPLICATION NUMBER: 60/235,834  
36 ; PRIOR FILING DATE: 2000-09-27  
37 ; PRIOR APPLICATION NUMBER: 60/234,274  
38 ; PRIOR FILING DATE: 2000-09-21  
39 ; PRIOR APPLICATION NUMBER: 60/234,223  
40 ; PRIOR FILING DATE: 2000-09-21  
41 ; PRIOR APPLICATION NUMBER: 60/228,924  
42 ; PRIOR FILING DATE: 2000-08-30  
43 ; PRIOR APPLICATION NUMBER: 60/224,518  
44 ; PRIOR FILING DATE: 2000-08-14  
45 ; PRIOR APPLICATION NUMBER: 60/236,369  
46 ; PRIOR FILING DATE: 2000-09-29  
47 ; PRIOR APPLICATION NUMBER: 60/224,519  
48 ; PRIOR FILING DATE: 2000-08-14  
49 ; PRIOR APPLICATION NUMBER: 60/220,964  
50 ; PRIOR FILING DATE: 2000-07-26  
51 ; PRIOR APPLICATION NUMBER: 60/241,809  
52 ; PRIOR FILING DATE: 2000-10-20  
53 ; PRIOR APPLICATION NUMBER: 60/249,299  
54 ; PRIOR FILING DATE: 2000-11-17  
55 ; PRIOR APPLICATION NUMBER: 60/236,327  
56 ; PRIOR FILING DATE: 2000-09-29  
57 ; PRIOR APPLICATION NUMBER: 60/241,785  
58 ; PRIOR FILING DATE: 2000-10-20  
59 ; PRIOR APPLICATION NUMBER: 60/244,617  
60 ; PRIOR FILING DATE: 2000-11-01  
61 ; PRIOR APPLICATION NUMBER: 60/225,268  
62 ; PRIOR FILING DATE: 2000-08-14  
63 ; PRIOR APPLICATION NUMBER: 60/236,368  
64 ; PRIOR FILING DATE: 2000-09-29  
65 ; PRIOR APPLICATION NUMBER: 60/251,856  
66 ; PRIOR FILING DATE: 2000-12-08  
67 ; PRIOR APPLICATION NUMBER: 60/251,868  
68 ; PRIOR FILING DATE: 2000-12-08  
69 ; PRIOR APPLICATION NUMBER: 60/229,344  
70 ; PRIOR FILING DATE: 2000-09-01  
71 ; PRIOR APPLICATION NUMBER: 60/234,997  
72 ; PRIOR FILING DATE: 2000-09-25  
73 ; PRIOR APPLICATION NUMBER: 60/229,343  
74 ; PRIOR FILING DATE: 2000-09-01  
75 ; PRIOR APPLICATION NUMBER: 60/229,345  
76 ; PRIOR FILING DATE: 2000-09-01  
77 ; PRIOR APPLICATION NUMBER: 60/229,287  
78 ; PRIOR FILING DATE: 2000-09-01  
79 ; PRIOR APPLICATION NUMBER: 60/229,513  
80 ; PRIOR FILING DATE: 2000-09-05  
81 ; PRIOR APPLICATION NUMBER: 60/231,413  
82 ; PRIOR FILING DATE: 2000-09-08  
83 ; PRIOR APPLICATION NUMBER: 60/229,509  
84 ; PRIOR FILING DATE: 2000-09-05  
85 ; PRIOR APPLICATION NUMBER: 60/236,367  
86 ; PRIOR FILING DATE: 2000-09-29  
87 ; PRIOR APPLICATION NUMBER: 60/237,039  
88 ; PRIOR FILING DATE: 2000-10-02  
89 ; PRIOR APPLICATION NUMBER: 60/237,038  
90 ; PRIOR FILING DATE: 2000-10-02  
91 ; PRIOR APPLICATION NUMBER: 60/236,370  
92 ; PRIOR FILING DATE: 2000-09-29  
93 ; PRIOR APPLICATION NUMBER: 60/236,802  
94 ; PRIOR FILING DATE: 2000-10-02  
95 ; PRIOR APPLICATION NUMBER: 60/237,037  
96 ; PRIOR FILING DATE: 2000-10-02  
97 ; PRIOR APPLICATION NUMBER: 60/237,040  
98 ; PRIOR FILING DATE: 2000-10-02  
99 ; PRIOR APPLICATION NUMBER: 60/240,960  
100 ; PRIOR FILING DATE: 2000-10-20  
101 ; PRIOR APPLICATION NUMBER: 60/239,935  
102 ; PRIOR FILING DATE: 2000-10-13  
103 ; PRIOR APPLICATION NUMBER: 60/239,937  
104 ; PRIOR FILING DATE: 2000-10-13  
105 ; PRIOR APPLICATION NUMBER: 60/241,787  
106 ; PRIOR FILING DATE: 2000-10-20  
107 ; PRIOR APPLICATION NUMBER: 60/246,474  
108 ; PRIOR FILING DATE: 2000-11-08  
109 ; PRIOR APPLICATION NUMBER: 60/246,532  
110 ; PRIOR FILING DATE: 2000-11-08  
111 ; PRIOR APPLICATION NUMBER: 60/249,216  
112 ; PRIOR FILING DATE: 2000-11-17  
113 ; PRIOR APPLICATION NUMBER: 60/249,210  
114 ; PRIOR FILING DATE: 2000-11-17  
115 ; PRIOR APPLICATION NUMBER: 60/226,681  
116 ; PRIOR FILING DATE: 2000-08-22  
117 ; PRIOR APPLICATION NUMBER: 60/225,759  
118 ; PRIOR FILING DATE: 2000-08-14  
119 ; PRIOR APPLICATION NUMBER: 60/225,213  
120 ; PRIOR FILING DATE: 2000-08-14  
121 ; PRIOR APPLICATION NUMBER: 60/227,182  
122 ; PRIOR FILING DATE: 2000-08-22  
123 ; PRIOR APPLICATION NUMBER: 60/225,214  
124 ; PRIOR FILING DATE: 2000-08-14  
125 ; PRIOR APPLICATION NUMBER: 60/235,836  
126 ; PRIOR FILING DATE: 2000-09-27  
127 ; PRIOR APPLICATION NUMBER: 60/230,438  
128 ; PRIOR FILING DATE: 2000-09-06  
129 ; PRIOR APPLICATION NUMBER: 60/215,135  
130 ; PRIOR FILING DATE: 2000-06-30  
131 ; PRIOR APPLICATION NUMBER: 60/225,266  
132 ; PRIOR FILING DATE: 2000-08-14  
133 ; PRIOR APPLICATION NUMBER: 60/249,218  
134 ; PRIOR FILING DATE: 2000-11-17  
135 ; PRIOR APPLICATION NUMBER: 60/249,208  
136 ; PRIOR FILING DATE: 2000-11-17  
137 ; PRIOR APPLICATION NUMBER: 60/249,213  
138 ; PRIOR FILING DATE: 2000-11-17  
139 ; PRIOR APPLICATION NUMBER: 60/249,212  
140 ; PRIOR FILING DATE: 2000-11-17  
141 ; PRIOR APPLICATION NUMBER: 60/249,207  
142 ; PRIOR FILING DATE: 2000-11-17  
143 ; PRIOR APPLICATION NUMBER: 60/249,245  
144 ; PRIOR FILING DATE: 2000-11-17  
145 ; PRIOR APPLICATION NUMBER: 60/249,244  
146 ; PRIOR FILING DATE: 2000-11-17

PRIOR APPLICATION NUMBER: 60/249,217  
PRIOR FILING DATE: 2000-11-17  
PRIOR APPLICATION NUMBER: 60/249,211  
PRIOR FILING DATE: 2000-11-17  
PRIOR APPLICATION NUMBER: 60/249,215  
PRIOR FILING DATE: 2000-11-17  
PRIOR APPLICATION NUMBER: 60/249,264  
PRIOR FILING DATE: 2000-11-17  
PRIOR APPLICATION NUMBER: 60/249,214  
PRIOR FILING DATE: 2000-11-17  
PRIOR APPLICATION NUMBER: 60/249,297  
PRIOR FILING DATE: 2000-11-17  
PRIOR APPLICATION NUMBER: 60/232,400  
PRIOR FILING DATE: 2000-09-14  
PRIOR APPLICATION NUMBER: 60/231,242  
PRIOR FILING DATE: 2000-09-08  
PRIOR APPLICATION NUMBER: 60/232,081  
PRIOR FILING DATE: 2000-09-08  
PRIOR APPLICATION NUMBER: 60/232,080  
PRIOR FILING DATE: 2000-09-08  
PRIOR APPLICATION NUMBER: 60/231,414  
PRIOR FILING DATE: 2000-09-08  
PRIOR APPLICATION NUMBER: 60/231,244  
PRIOR FILING DATE: 2000-09-08  
PRIOR APPLICATION NUMBER: 60/233,064  
PRIOR FILING DATE: 2000-09-14  
PRIOR APPLICATION NUMBER: 60/233,063  
PRIOR FILING DATE: 2000-09-14  
PRIOR APPLICATION NUMBER: 60/232,397  
PRIOR FILING DATE: 2000-09-14  
PRIOR APPLICATION NUMBER: 60/232,399  
PRIOR FILING DATE: 2000-09-14  
PRIOR APPLICATION NUMBER: 60/232,401  
PRIOR FILING DATE: 2000-09-14  
PRIOR APPLICATION NUMBER: 60/241,808  
PRIOR FILING DATE: 2000-10-20  
PRIOR APPLICATION NUMBER: 60/241,826  
PRIOR FILING DATE: 2000-10-20  
PRIOR APPLICATION NUMBER: 60/241,786  
PRIOR FILING DATE: 2000-10-20  
PRIOR APPLICATION NUMBER: 60/241,221  
PRIOR FILING DATE: 2000-10-20  
PRIOR APPLICATION NUMBER: 60/246,475  
PRIOR FILING DATE: 2000-11-08  
PRIOR APPLICATION NUMBER: 60/231,243  
PRIOR FILING DATE: 2000-09-08

Query Match 32.0%; Score 31; DB 14; Length 50;  
Best Local Similarity 60.0%; Pred. No. 1e+03;  
Matches 1; Mismatches 3; Indels 0; Gaps 0;

QY 2 PNLNSKIAF 11  
DB 36 PPHVNWKTAF 45

## RESULT 42

US-10-212-872-359  
Sequence 359, Application US/10212872  
Publication No. US20030215893A1  
GENERAL INFORMATION:  
APPLICANT: Rosen et al.  
TITLE OF INVENTION: Nucleic Acids, Proteins, and Antibodies  
FILE REFERENCE: PC008C2  
CURRENT APPLICATION NUMBER: US/10/212,872  
CURRENT FILING DATE: 2002-08-07  
Prior application removed - See File Wrapper or Palm  
NUMBER OF SEQ ID NOS: 1198  
SOFTWARE: PatentIn Ver. 2.0  
SEQ ID NO 359  
LENGTH: 50  
TYPE: PRT  
ORGANISM: Homo sapiens

US-10-212-872-359

Query Match 32.0%; Score 31; DB 14; Length 50;  
Best Local Similarity 60.0%; Pred. No. 1e+03;  
Matches 6; Conservative 1; Mismatches 3; Indels 0; Gaps 0;

QY 2 PNLNSKIAF 11  
DB 36 PPHVNWKTAF 45

## RESULT 43

US-10-424-599-181551  
Sequence 181551, Application US/10424599  
Publication No. US20040031072A1  
GENERAL INFORMATION:  
APPLICANT: Ia Rosa Thomas J  
APPLICANT: Kovalic David K  
APPLICANT: Zhou Yihua  
APPLICANT: Cao Yongwei  
TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With  
FILE REFERENCE: 38-21(53223)B  
CURRENT APPLICATION NUMBER: US/10/424,599  
CURRENT FILING DATE: 2003-04-28  
NUMBER OF SEQ ID NOS: 285684  
SEQ ID NO 181551  
LENGTH: 50  
TYPE: PRT  
ORGANISM: Glycine max  
FEATURE:  
OTHER INFORMATION: Clone ID: PAT\_MRT3847\_134955C.1.pap  
US-10-424-599-181551

Query Match 32.0%; Score 31; DB 15; Length 50;  
Best Local Similarity 70.0%; Pred. No. 1e+03;  
Matches 7; Conservative 0; Mismatches 3; Indels 0; Gaps 0;

QY 6 NSKIAFKIVS 15  
DB 11 NKKICFPVVS 20

## RESULT 44

US-10-767-701-52857  
Sequence 52857, Application US/10767701  
Publication No. US20040172684A1  
GENERAL INFORMATION:  
APPLICANT: Kovalic, David K.  
APPLICANT: Zhou, Yihua  
APPLICANT: Cao, Yongwei  
TITLE OF INVENTION: Nucleic Acid Molecules and Other Molecules Associated With  
FILE REFERENCE: 38-21(53535)B  
CURRENT APPLICATION NUMBER: US/10/767,701  
CURRENT FILING DATE: 2004-01-29  
NUMBER OF SEQ ID NOS: 63128  
SEQ ID NO 52857  
LENGTH: 50  
TYPE: PRT  
ORGANISM: Sorghum bicolor  
FEATURE:  
OTHER INFORMATION: Clone ID: 13152278.pap  
US-10-767-701-52857

Query Match 32.0%; Score 31; DB 16; Length 50;  
Best Local Similarity 83.3%; Pred. No. 1e+03;  
Matches 5; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 1 EPNHLN 6  
DB 23 EPNHLH 28

## RESULT 45

US-10-153-344-10  
 ; Sequence 10, Application US/10153344  
 ; Publication No. US20030004124A1  
 ; GENERAL INFORMATION:  
 ; APPLICANT: ROTHMAN, JOEL  
 ; APPLICANT: BLOSS, TIM  
 ; APPLICANT: WITZE, ERIC  
 ; TITLE OF INVENTION: BTF3: AN INHIBITOR OF APOPTOSIS  
 ; FILE REFERENCE: 407T-300410US  
 ; CURRENT APPLICATION NUMBER: US/10/153,344  
 ; CURRENT FILING DATE: 2002-08-27  
 ; PRIOR APPLICATION NUMBER: US 60/292,559  
 ; PRIOR FILING DATE: 2001-05-21  
 ; NUMBER OF SEQ ID NOS: 35  
 ; SOFTWARE: PatentIn version 3.0  
 ; SEQ ID NO 10  
 ; LENGTH: 19  
 ; TYPE: PRT  
 ; ORGANISM: Caenorhabditis elegans  
 US-10-153-344-10

Query Match 31.4%; Score 30.5; DB 14; Length 19;  
 Best Local Similarity 47.1%; Pred. No. 4.3e+02;  
 Matches 8; Conservative 4; Mismatches 2; Indels 3; Gaps 1;

## QY 3 NHLNSKIAPKIVSQBPA 19

Db 2 DHLRAK--KILSREDA 15  
 :||:| ||:|:|

## RESULT 46

US-10-424-599-212560  
 ; Sequence 212560, Application US/10424599  
 ; Publication No. US20040031072A1  
 ; GENERAL INFORMATION:  
 ; APPLICANT: La Rosa Thomas J  
 ; APPLICANT: Kovalic David K  
 ; APPLICANT: Zhou Yihua  
 ; APPLICANT: Cao Yongwei  
 ; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With  
 ; FILE REFERENCE: 38-21(53223)B  
 ; CURRENT APPLICATION NUMBER: US/10/424,599  
 ; CURRENT FILING DATE: 2003-04-28  
 ; NUMBER OF SEQ ID NOS: 285684  
 ; SEQ ID NO 212560  
 ; LENGTH: 37  
 ; TYPE: PRT  
 ; ORGANISM: Glycine max  
 ; FEATURE:  
 ; OTHER INFORMATION: Clone ID: PAT\_MRT3847\_33969C.1.pep  
 US-10-424-599-212560

Query Match 31.4%; Score 30.5; DB 15; Length 37;  
 Best Local Similarity 50.0%; Pred. No. 9.1e+02;  
 Matches 7; Conservative 3; Mismatches 1; Indels 3; Gaps 1;

## QY 2 PNLNSKIAPKIVS 15

Db 3 PNLNS--YKVLS 13  
 ||||| :|:|

## RESULT 47

US-10-282-122A-71852  
 ; Sequence 71852, Application US/10282122A  
 ; Publication No. US20040029129A1  
 ; GENERAL INFORMATION:  
 ; APPLICANT: Wang, Liangsu  
 ; APPLICANT: Zamudio, Carlos  
 ; APPLICANT: Malone, Cheryl  
 ; APPLICANT: Haselbeck, Robert

; APPLICANT: Ohlsen, Kari  
 ; APPLICANT: Zyskind, Judith  
 ; APPLICANT: Wall, Daniel  
 ; APPLICANT: Trawick, John  
 ; APPLICANT: Carr, Grant  
 ; APPLICANT: Yamamoto, Robert  
 ; APPLICANT: Forsyth, R.  
 ; APPLICANT: Xu, H.  
 ; TITLE OF INVENTION: Identification of Essential Genes in Microorganisms  
 ; FILE REFERENCE: ELITRA.034A  
 ; CURRENT APPLICATION NUMBER: US/10/282,122A  
 ; CURRENT FILING DATE: 2003-02-20  
 ; PRIOR APPLICATION NUMBER: 60/191,078  
 ; PRIOR FILING DATE: 2000-03-21  
 ; PRIOR APPLICATION NUMBER: 60/206,848  
 ; PRIOR FILING DATE: 2000-05-23  
 ; PRIOR APPLICATION NUMBER: 60/207,727  
 ; PRIOR FILING DATE: 2000-05-26  
 ; PRIOR APPLICATION NUMBER: 60/230,335  
 ; PRIOR FILING DATE: 2000-09-06  
 ; PRIOR APPLICATION NUMBER: 60/230,347  
 ; PRIOR FILING DATE: 2000-09-09  
 ; PRIOR APPLICATION NUMBER: 60/242,578  
 ; PRIOR FILING DATE: 2000-10-23  
 ; PRIOR APPLICATION NUMBER: 60/253,625  
 ; PRIOR FILING DATE: 2000-11-27  
 ; PRIOR APPLICATION NUMBER: 60/257,931  
 ; PRIOR FILING DATE: 2000-12-22  
 ; PRIOR APPLICATION NUMBER: 60/267,636  
 ; PRIOR FILING DATE: 2001-02-09  
 ; PRIOR APPLICATION NUMBER: 60/269,308  
 ; PRIOR FILING DATE: 2001-02-16  
 ; Remaining Prior Application data removed - See File Wrapper or PALM.  
 ; NUMBER OF SEQ ID NOS: 78614  
 ; SOFTWARE: PatentIn version 3.1  
 ; SEQ ID NO 71852  
 ; LENGTH: 40  
 ; TYPE: PRT  
 ; ORGANISM: Staphylococcus haemolyticus  
 ; FEATURE:  
 ; NAME/KEY: MISC\_FEATURE  
 ; LOCATION: (31)..(31)  
 ; OTHER INFORMATION: X=any amino acid  
 US-10-282-122A-71852  
 Query Match 31.4%; Score 30.5; DB 15; Length 40;  
 Best Local Similarity 37.5%; Pred. No. 9.9e+02;  
 Matches 6; Conservative 6; Mismatches 3; Indels 1; Gaps 1;  
 QY 1 EPNHLSKIAPKIVSQ 16  
 :|||:|:|:|  
 Db 15 DPH-NSKLVTLINK 29  
 :|||:|:|:|  
 RESULT 48  
 US-10-424-599-144936  
 ; Sequence 144936, Application US/10424599  
 ; Publication No. US20040031072A1  
 ; GENERAL INFORMATION:  
 ; APPLICANT: La Rosa Thomas J  
 ; APPLICANT: Kovalic David K  
 ; APPLICANT: Zhou Yihua  
 ; APPLICANT: Cao Yongwei  
 ; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With  
 ; FILE REFERENCE: 38-21(53223)B  
 ; CURRENT APPLICATION NUMBER: US/10/424,599  
 ; CURRENT FILING DATE: 2003-04-28  
 ; NUMBER OF SEQ ID NOS: 285684  
 ; SEQ ID NO 144936  
 ; LENGTH: 40  
 ; TYPE: PRT  
 ; ORGANISM: Glycine max

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;
; FEATURE:
; OTHER INFORMATION: Clone ID: PAT_MRT3847_101896C.1.pep
; US-10-424-599-144936

Query Match      31.4%; Score 30.5; DB 15; Length 40;
Best Local Similarity 35.3%; Pred. No. 9.9e+02;
Matches 6; Conservative 5; Mismatches 5; Indels 1; Gaps 1;

QY 2 PNHLSKIAPKIVSQEP 18
   |:|:|:|:|:|:|
Db 25 PDH-GTKVSAKILQHP 40

RESULT 49
US-10-139-794-103
; Sequence 103, Application US/10139794
; Publication No. US20030232421A1
; GENERAL INFORMATION:
; APPLICANT: HYBRIGENICS, LYNX THERAPEUTICS INC.
; TITLE OF INVENTION: Protein-Protein Interactions In Adipocyte Cells (3)
; FILE REFERENCE: B4883A
; CURRENT APPLICATION NUMBER: US/10/139,794
; PRIOR APPLICATION NUMBER: US 60/288,885
; PRIOR FILING DATE: 2002-05-06
; PRIOR FILING DATE: 2001-05-04
; NUMBER OF SEQ ID NOS: 2930
; SOFTWARE: PatentIn version 3.1
; SEQ ID NO 103
; LENGTH: 25
; TYPE: PRT
; ORGANISM: Homo Sapiens
; FEATURE:
; OTHER INFORMATION: Translation of SEQ ID NO:102
US-10-139-794-103

Query Match      30.9%; Score 30; DB 14; Length 25;
Best Local Similarity 56.2%; Pred. No. 7.1e+02;
Matches 9; Conservative 1; Mismatches 4; Indels 2; Gaps 1;

QY 1 EPNHL--NSKIAFKIV 14
   |||:|:|:|:|
Db 3 EPKHLLEGITASKIV 18

RESULT 50
US-09-864-761-48976
; Sequence 48976, Application US/09864761
; Patent No. US20020048763A1
; GENERAL INFORMATION:
; APPLICANT: Penn, Sharon G.
; APPLICANT: Rank, David R.
; APPLICANT: Hanzel, David K.
; APPLICANT: Chen, Wensheng
; TITLE OF INVENTION: HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES USEFUL FOR
; TITLE OF INVENTION: GENE EXPRESSION ANALYSIS BY MICROARRAY
; FILE REFERENCE: Aecmica-X-1
; CURRENT APPLICATION NUMBER: US/09/864,761
; CURRENT FILING DATE: 2001-05-23
; PRIOR APPLICATION NUMBER: US 60/180,312
; PRIOR FILING DATE: 2000-02-04
; PRIOR APPLICATION NUMBER: US 60/207,456
; PRIOR FILING DATE: 2000-05-26
; PRIOR APPLICATION NUMBER: US 09/632,366
; PRIOR FILING DATE: 2000-08-03
; PRIOR APPLICATION NUMBER: GB 24263.6
; PRIOR FILING DATE: 2000-10-04
; PRIOR APPLICATION NUMBER: US 60/236,359
; PRIOR FILING DATE: 2000-09-27
; PRIOR APPLICATION NUMBER: PCT/US01/00666
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00667
; PRIOR FILING DATE: 2001-01-30
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;
; PRIOR APPLICATION NUMBER: PCT/US01/00664
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00669
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00665
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00668
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00663
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00662
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00661
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: PCT/US01/00670
; PRIOR FILING DATE: 2001-01-30
; PRIOR APPLICATION NUMBER: US 60/234,687
; PRIOR FILING DATE: 2000-09-21
; PRIOR APPLICATION NUMBER: US 09/608,408
; PRIOR FILING DATE: 2000-06-30
; PRIOR APPLICATION NUMBER: US 09/774,203
; PRIOR FILING DATE: 2001-01-29
; NUMBER OF SEQ ID NOS: 49117
; SOFTWARE: Annomax Sequence Listing Engine vers. 1.1
; SEQ ID NO 48976
; LENGTH: 26
; TYPE: PRT
; ORGANISM: Homo sapiens
; FEATURE:
; OTHER INFORMATION: MAP TO AC011402.1
; OTHER INFORMATION: EXPRESSED IN HEL100, SIGNAL = 1.2
; OTHER INFORMATION: EXPRESSED IN HELA, SIGNAL = 1.9
US-09-864-761-48976

Query Match      30.9%; Score 30; DB 9; Length 26;
Best Local Similarity 46.2%; Pred. No. 7.4e+02;
Matches 6; Conservative 2; Mismatches 5; Indels 0; Gaps 0;

QY 1 EPNHLSKIAPKI 13
   |:|:|:|:|
Db 1 EKHHLDQMIKFSI 13

RESULT 51
US-09-933-767-898
; Sequence 898, Application US/09933767
; Publication No. US20030181692A1
; GENERAL INFORMATION:
; APPLICANT: Ni et al.
; TITLE OF INVENTION: 207 Human Secreted Proteins
; FILE REFERENCE: P2007E2
; CURRENT APPLICATION NUMBER: US/09/933,767
; CURRENT FILING DATE: 2001-08-22
; PRIOR APPLICATION NUMBER: PCT/US01/05614
; PRIOR FILING DATE: 2001-02-21
; PRIOR APPLICATION NUMBER: 60/184,836
; PRIOR FILING DATE: 2000-02-24
; PRIOR APPLICATION NUMBER: 60/193,170
; PRIOR FILING DATE: 2000-03-29
; PRIOR APPLICATION NUMBER: 09/205,258
; PRIOR FILING DATE: 1998-12-04
; PRIOR APPLICATION NUMBER: PCT/US98/11422
; PRIOR FILING DATE: 1998-06-04
; PRIOR APPLICATION NUMBER: 60/048,885
; PRIOR FILING DATE: 1997-06-06
; PRIOR APPLICATION NUMBER: 60/049,375
; PRIOR FILING DATE: 1997-06-06
; PRIOR APPLICATION NUMBER: 60/048,881
; PRIOR FILING DATE: 1997-06-06
; PRIOR APPLICATION NUMBER: 60/048,880
; PRIOR FILING DATE: 1997-06-06
; PRIOR APPLICATION NUMBER: 60/048,896
; PRIOR FILING DATE: 1997-06-06
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; PRIOR APPLICATION NUMBER: 60/085,923
; PRIOR FILING DATE: 1998-05-18
; PRIOR APPLICATION NUMBER: 60/085,922
; PRIOR FILING DATE: 1998-05-18
; PRIOR APPLICATION NUMBER: 60/092,921
; PRIOR FILING DATE: 1998-07-15
; PRIOR APPLICATION NUMBER: 60/094,657
; PRIOR FILING DATE: 1998-07-30
; NUMBER OF SEQ ID NOS: 1245
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 899
; LENGTH: 27
; TYPE: PRT
; ORGANISM: Homo sapiens
US-09-933-767-899

Query Match          30.9%; Score 30; DB 10; Length 27;
Best Local Similarity 37.5%; Pred. No. 7.7e+02;
Matches 6; Conservative 4; Mismatches 6; Indels

QY 2 PNHLNSKIAFKIVSQE 17
Db 6 PSANNQRFAPSPLEE 21

RESULT 53
US-10-004-860-898
; Sequence 898, Application US/10004860
; Publication No. US20030065160A1
; GENERAL INFORMATION:
; APPLICANT: Young et al.
; TITLE OF INVENTION: 207 Human Secreted Proteins
; FILE REFERENCE: PZ007P1
; CURRENT APPLICATION NUMBER: US/10/004,860
; PRIOR FILING DATE: 2001-12-07
; Prior Application removed - See File Wrapper or Palm
; NUMBER OF SEQ ID NOS: 1227
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 898
; LENGTH: 27
; TYPE: PRT
; ORGANISM: Homo sapiens
US-10-004-860-898

Query Match          30.9%; Score 30; DB 14; Length 27;
Best Local Similarity 37.5%; Pred. No. 7.7e+02;
Matches 6; Conservative 4; Mismatches 6; Indels

Qy 2 PNHLNSKIAFKIVSQE 17
Db 6 PSANNQRFAPSPLEE 21

RESULT 54
US-10-004-860-899
; Sequence 899, Application US/10004860
; Publication No. US20030065160A1
; GENERAL INFORMATION:
; APPLICANT: Young et al.
; TITLE OF INVENTION: 207 Human Secreted Proteins
; FILE REFERENCE: PZ007P1
; CURRENT APPLICATION NUMBER: US/10/004,860
; CURRENT FILING DATE: 2001-12-07
; Prior Application removed - See File Wrapper or Palm
; NUMBER OF SEQ ID NOS: 1227
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 899
; LENGTH: 27
; TYPE: PRT
; ORGANISM: Homo sapiens
US-10-004-860-899

Query Match          30.9%; Score 30; DB 14; Length 27;

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Best Local Similarity 37.5%; Pred. No. 7.7e+02;
Matches 6; Conservative 4; Mismatches 6; Indels 0; Gaps 0;

QY 2 PNHLSKIAFKIVSQE 17
   | : | : | : | : |
Db 6 PSANNQRFAPSLSEE 21

RESULT 55
US-10-023-282-898
; Sequence 898, Application US/10023282
; Publication No. US20030092893A1
; GENERAL INFORMATION:
; APPLICANT: Young et al.
; TITLE OF INVENTION: 207 Human Secreted Proteins
; FILE REFERENCE: PZ007P1
; CURRENT APPLICATION NUMBER: US/10/023,282
; CURRENT FILING DATE: 2001-12-20
; EARLIER APPLICATION NUMBER: 09/205,258
; EARLIER FILING DATE: 1998-12-04
; EARLIER APPLICATION NUMBER: 60/048,885
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/049,375
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/048,881
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/048,880
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/048,896
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/049,020
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/048,876
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/049,374
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/048,895
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/048,892
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/048,900
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/048,901
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/048,892
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/048,915
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/049,019
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/048,970
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/048,972
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/048,916
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/049,373
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/048,875
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/049,374

Query Match 30.9%; Score 30; DB 14; Length 27;
Best Local Similarity 37.5%; Pred. No. 7.7e+02;
Matches 6; Conservative 4; Mismatches 6; Indels 0; Gaps 0;

QY 2 PNHLSKIAFKIVSQE 17
   | : | : | : | : |
Db 6 PSANNQRFAPSLSEE 21

RESULT 56
US-10-023-282-899
; Sequence 899, Application US/10023282
; Publication No. US20030092893A1
; GENERAL INFORMATION:
; APPLICANT: Young et al.
; TITLE OF INVENTION: 207 Human Secreted Proteins
; FILE REFERENCE: PZ007P1
; CURRENT APPLICATION NUMBER: US/10/023,282
; CURRENT FILING DATE: 2001-12-20
; EARLIER APPLICATION NUMBER: 09/205,258
; EARLIER FILING DATE: 1998-12-04
; EARLIER APPLICATION NUMBER: PCT/US98/11422
; EARLIER FILING DATE: 1998-06-04
; EARLIER APPLICATION NUMBER: 60/048,885
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/049,375
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/048,881
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/048,880
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/048,896
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/049,020
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/048,876
; EARLIER FILING DATE: 1997-06-06
; EARLIER APPLICATION NUMBER: 60/048,895
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Query Match	30.9%;	Score 30;	DB 14;	Length 27;
Best Local Similarity	37.5%;	Pred. No. 7.7e+02.		

Db 13 HLNINFIKLDQQ 25

RESULT 59

US-10-424-599-207820

; Sequence 207820, Application US/10424599

; Publication No. US20040031072A1

; GENERAL INFORMATION:

; APPLICANT: La Rosa Thomas J

; APPLICANT: Kovalic David K

; APPLICANT: Zhou Yihua

; APPLICANT: Cao Yongwei

; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With

; TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement

; FILE REFERENCE: 38-21(53223)B

; CURRENT APPLICATION NUMBER: US/10/424,599

; CURRENT FILING DATE: 2003-04-28

; NUMBER OF SEQ ID NOS: 285684

; SEQ ID NO 207820

; LENGTH: 30

; TYPE: PRT

; ORGANISM: Glycine max

; FEATURE:

; OTHER INFORMATION: Clone ID: PAT\_MRT3847\_29688C.1.pap

US-10-424-599-207820

Query Match 30.9%; Score 30; DB 15; Length 30;

Best Local Similarity 33.3%; Pred. No. 8.7e+02;

Matches 5; Conservative 3; Mismatches 7; Indels 0; Gaps 0;

QY 2 PNHLNSKIAPKIVSQ 16

Db 4 PTHIKVKLVKHPQ 18

RESULT 60

US-10-296-734-654

; Sequence 654, Application US/10296734

; Publication No. US20040054137A1

; GENERAL INFORMATION:

; APPLICANT: Thompson, Scott A

; APPLICANT: Ramshaw, Ian A

; TITLE OF INVENTION: Synthetic molecules and uses therefor

; FILE REFERENCE: Savine

; CURRENT APPLICATION NUMBER: US/10/296,734

; CURRENT FILING DATE: 2003-08-04

; PRIOR APPLICATION NUMBER: AU PQ7761/00

; PRIOR FILING DATE: 2000-05-26

; NUMBER OF SEQ ID NOS: 1507

; SOFTWARE: PatentIn version 3.2

; SEQ ID NO 654

; LENGTH: 30

; TYPE: PRT

; ORGANISM: Artificial

; FEATURE:

; OTHER INFORMATION: HepC 1a segment 124

US-10-296-734-654

Query Match 30.9%; Score 30; DB 15; Length 30;

Best Local Similarity 56.7%; Pred. No. 8.7e+02;

Matches 6; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

QY 9 IAPKIVSQE 17

Db 21 VAFKIMSGE 29

RESULT 61

US-10-296-734-656

; Sequence 656, Application US/10296734

; Publication No. US20040054137A1

; GENERAL INFORMATION:

; APPLICANT: Thompson, Scott A

; APPLICANT: Ramshaw, Ian A

; TITLE OF INVENTION: Synthetic molecules and uses therefor

; FILE REFERENCE: Savine

; CURRENT APPLICATION NUMBER: US/10/296,734

; CURRENT FILING DATE: 2003-08-04

; PRIOR APPLICATION NUMBER: AU PQ7761/00

; PRIOR FILING DATE: 2000-05-26

; NUMBER OF SEQ ID NOS: 1507

; SOFTWARE: PatentIn version 3.2

; SEQ ID NO 656

; LENGTH: 30

; TYPE: PRT

; ORGANISM: Artificial

; FEATURE:

; OTHER INFORMATION: HepC 1a segment 125

US-10-296-734-656

Query Match 30.9%; Score 30; DB 15; Length 30;

Best Local Similarity 66.7%; Pred. No. 8.7e+02;

Matches 6; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

QY 9 IAPKIVSQE 17

Db 6 VAFKIMSGE 14

RESULT 62

US-10-050-704-215

; Sequence 215, Application US/10050704

; Publication No. US20030050442A1

; GENERAL INFORMATION:

; APPLICANT: Ruben et al.

; TITLE OF INVENTION: 62 Human Secreted Proteins

; FILE REFERENCE: PZ039P1

; CURRENT APPLICATION NUMBER: US/10/050,704

; CURRENT FILING DATE: 2002-01-18

; PRIOR APPLICATION NUMBER: 09/684,524

; PRIOR FILING DATE: 2000-10-10

; PRIOR APPLICATION NUMBER: PCT/US00/08979

; PRIOR FILING DATE: 2000-04-06

; PRIOR APPLICATION NUMBER: 60/128,693

; PRIOR FILING DATE: 1999-04-09

; PRIOR APPLICATION NUMBER: 60/130,991

; PRIOR FILING DATE: 1999-04-26

; NUMBER OF SEQ ID NOS: 344

; SOFTWARE: PatentIn Ver. 2.0

; SEQ ID NO 215

; LENGTH: 31

; TYPE: PRT

; ORGANISM: Homo sapiens

US-10-050-704-215

Query Match 30.9%; Score 30; DB 14; Length 31;

Best Local Similarity 33.3%; Pred. No. 9e+02;

Matches 5; Conservative 3; Mismatches 7; Indels 0; Gaps 0;

QY 2 PNHLNSKIAPKIVSQ 16

Db 5 PNNIRHKFGSNVVDQ 19

RESULT 63

US-10-798-512-215

; Sequence 215, Application US/10798512

; Publication No. US20040152164A1

; GENERAL INFORMATION:

; APPLICANT: Ruben et al.

; TITLE OF INVENTION: 62 Human Secreted Proteins

; FILE REFERENCE: PZ039P1

; CURRENT APPLICATION NUMBER: US/10/798,512

; CURRENT FILING DATE: 2004-03-12

; PRIOR APPLICATION NUMBER: US/03/684,524

; PRIOR FILING DATE: 2000-10-10

Db 13 HLNINFIKLDQQ 25

RESULT 59

US-10-424-599-207820

; Sequence 207820, Application US/10424599

; Publication No. US20040031072A1

; GENERAL INFORMATION:

; APPLICANT: La Rosa Thomas J

; APPLICANT: Kovalic David K

; APPLICANT: Zhou Yihua

; APPLICANT: Cao Yongwei

; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With

; TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement

; FILE REFERENCE: 38-21(53223)B

; CURRENT APPLICATION NUMBER: US/10/424,599

; CURRENT FILING DATE: 2003-04-28

; NUMBER OF SEQ ID NOS: 285684

; SEQ ID NO 207820

; LENGTH: 30

; TYPE: PRT

; ORGANISM: Glycine max

; FEATURE:

; OTHER INFORMATION: Clone ID: PAT\_MRT3847\_29688C.1.pap

US-10-424-599-207820

Query Match 30.9%; Score 30; DB 15; Length 30;

Best Local Similarity 33.3%; Pred. No. 8.7e+02;

Matches 5; Conservative 3; Mismatches 7; Indels 0; Gaps 0;

QY 2 PNHLNSKIAPKIVSQ 16

Db 4 PTHIKVKLVKHPQ 18

RESULT 60

US-10-296-734-654

; Sequence 654, Application US/10296734

; Publication No. US20040054137A1

; GENERAL INFORMATION:

; APPLICANT: Thompson, Scott A

; APPLICANT: Ramshaw, Ian A

; TITLE OF INVENTION: Synthetic molecules and uses therefor

; FILE REFERENCE: Savine

; CURRENT APPLICATION NUMBER: US/10/296,734

; CURRENT FILING DATE: 2003-08-04

; PRIOR APPLICATION NUMBER: AU PQ7761/00

; PRIOR FILING DATE: 2000-05-26

; NUMBER OF SEQ ID NOS: 1507

; SOFTWARE: PatentIn version 3.2

; SEQ ID NO 654

; LENGTH: 30

; TYPE: PRT

; ORGANISM: Artificial

; FEATURE:

; OTHER INFORMATION: HepC 1a segment 124

US-10-296-734-654

Query Match 30.9%; Score 30; DB 15; Length 30;

Best Local Similarity 56.7%; Pred. No. 8.7e+02;

Matches 6; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

QY 9 IAPKIVSQE 17

Db 21 VAFKIMSGE 29

RESULT 61

US-10-296-734-656

; Sequence 656, Application US/10296734

; Publication No. US20040054137A1

; GENERAL INFORMATION:

; APPLICANT: Thompson, Scott A

; APPLICANT: Ramshaw, Ian A

; TITLE OF INVENTION: Synthetic molecules and uses therefor

; FILE REFERENCE: Savine

; CURRENT APPLICATION NUMBER: US/10/296,734

; CURRENT FILING DATE: 2003-08-04

; PRIOR APPLICATION NUMBER: AU PQ7761/00

; PRIOR FILING DATE: 2000-05-26

; NUMBER OF SEQ ID NOS: 1507

; SOFTWARE: PatentIn version 3.2

; SEQ ID NO 656

; LENGTH: 30

; TYPE: PRT

; ORGANISM: Artificial

; FEATURE:

; OTHER INFORMATION: HepC 1a segment 125

US-10-296-734-656

Query Match 30.9%; Score 30; DB 15; Length 30;

Best Local Similarity 66.7%; Pred. No. 8.7e+02;

Matches 6; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

QY 9 IAPKIVSQE 17

Db 6 VAFKIMSGE 14

RESULT 62

US-10-050-704-215

; Sequence 215, Application US/10050704

; Publication No. US20030050442A1

; GENERAL INFORMATION:

; APPLICANT: Ruben et al.

; TITLE OF INVENTION: 62 Human Secreted Proteins

; FILE REFERENCE: PZ039P1

; CURRENT APPLICATION NUMBER: US/10/050,704

; CURRENT FILING DATE: 2002-01-18

; PRIOR APPLICATION NUMBER: 09/684,524

; PRIOR FILING DATE: 2000-10-10

; PRIOR APPLICATION NUMBER: PCT/US00/08979

; PRIOR FILING DATE: 2000-04-06

; PRIOR APPLICATION NUMBER: 60/128,693

; PRIOR FILING DATE: 1999-04-09

; PRIOR APPLICATION NUMBER: 60/130,991

; PRIOR FILING DATE: 1999-04-26

; NUMBER OF SEQ ID NOS: 344

; SOFTWARE: PatentIn Ver. 2.0

; SEQ ID NO 215

; LENGTH: 31

; TYPE: PRT

; ORGANISM: Homo sapiens

US-10-050-704-215

Query Match 30.9%; Score 30; DB 14; Length 31;

Best Local Similarity 33.3%; Pred. No. 9e+02;

Matches 5; Conservative 3; Mismatches 7; Indels 0; Gaps 0;

QY 2 PNHLNSKIAPKIVSQ 16

Db 5 PNNIRHKFGSNVVDQ 19

RESULT 63

US-10-798-512-215

; Sequence 215, Application US/10798512

; Publication No. US20040152164A1

; GENERAL INFORMATION:

; APPLICANT: Ruben et al.

; TITLE OF INVENTION: 62 Human Secreted Proteins

; FILE REFERENCE: PZ039P1

; CURRENT APPLICATION NUMBER: US/10/798,512

; CURRENT FILING DATE: 2004-03-12

; PRIOR APPLICATION NUMBER: US/03/684,524

; PRIOR FILING DATE: 2000-10-10

;; PRIOR APPLICATION NUMBER: PCT/US00/08979  
;; PRIOR FILING DATE: 2000-04-06  
;; PRIOR APPLICATION NUMBER: 60/128,693  
;; PRIOR FILING DATE: 1999-04-09  
;; PRIOR APPLICATION NUMBER: 60/130,991  
;; PRIOR FILING DATE: 1999-04-26  
;; NUMBER OF SEQ ID NOS: 344  
;; SOFTWARE: PatentIn Ver. 2.0  
;; SEQ ID NO 215  
;; LENGTH: 31  
;; TYPE: PRT  
;; ORGANISM: Homo sapiens  
US-10-798-512-215

Query Match 30.9%; Score 30; DB 16; Length 31;  
Best Local Similarity 33.3%; Pred. No. 9e+02;  
Matches 5; Conservative 3; Mismatches 7; Indels 0; Gaps 0;

QY 2 PNHLSKIAFKIVSQ 16  
||:|:|:  
DB 5 PNIHRKFGSNVDQ 19

RESULT 64  
US-10-106-698-5968  
; Sequence 5968, Application US/10106698  
; Publication No. US20030109690A1  
; GENERAL INFORMATION:  
; APPLICANT: Ruben et al.  
; TITLE OF INVENTION: Colon and Colon Cancer Associated Polynucleotides and Polypeptide  
; FILE REFERENCE: PA005P1  
; CURRENT APPLICATION NUMBER: US/10/106,698  
; CURRENT FILING DATE: 2002-03-27  
; PRIOR APPLICATION NUMBER: PCT/US00/26524  
; PRIOR FILING DATE: 2000-09-28  
; PRIOR APPLICATION NUMBER: US 60/157,137  
; PRIOR FILING DATE: 1999-09-29  
; PRIOR APPLICATION NUMBER: US 60/163,280  
; PRIOR FILING DATE: 1999-11-03  
; NUMBER OF SEQ ID NOS: 8564  
; SOFTWARE: PatentIn Ver. 3.0  
; SEQ ID NO 5968  
; LENGTH: 32  
; TYPE: PRT  
; ORGANISM: Homo sapiens  
; NAME/KEY: MISC\_FEATURE  
; LOCATION: (20)  
; OTHER INFORMATION: Xaa equals any of the naturally occurring L-amino acids  
US-10-106-698-5968

Query Match 30.9%; Score 30; DB 14; Length 32;  
Best Local Similarity 77.8%; Pred. No. 9.3e+02;  
Matches 7; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

QY 8 KIAFKIVSQ 16  
||:|:|:  
DB 5 KIAWKIVIQ 13

RESULT 65  
US-10-300-083-13  
; Sequence 13, Application US/10300083  
; Publication No. US20030153502A1  
; GENERAL INFORMATION:  
; APPLICANT: REGENTS OF THE UNIVERSITY OF MINNESOTA  
; TITLE OF INVENTION: SYNTHETIC APPROACH TO DESIGNED CHEMICAL STRUCTURES  
; NUMBER OF SEQUENCES: 38  
; CORRESPONDENCE ADDRESS:  
; ADDRESSEE: MUEITING, RAASCH, GEBHARDT & SCHWAPPACH, P.A.  
; STREET: 119 No. US20030153502A1th Fourth Street, Suite 203  
; CITY: Minneapolis

;; STATE: Minnesota  
;; COUNTRY: U.S.A.  
;; ZIP: 55401  
;; COMPUTER READABLE FORM:  
;; MEDIUM TYPE: Floppy disk  
;; COMPUTER: IBM PC compatible  
;; OPERATING SYSTEM: PC-DOS/MS-DOS  
;; SOFTWARE: PatentIn Release #1.0, Version #1.30  
;; CURRENT APPLICATION DATA:  
;; APPLICATION NUMBER: US/10/300,083  
;; FILING DATE: 20-No. US20030153502A1-2002  
;; CLASSIFICATION: <Unknown>  
;; PRIOR APPLICATION DATA:  
;; APPLICATION NUMBER: US/09/194,296  
;; FILING DATE: 15-Oct-1999  
;; APPLICATION NUMBER: US 08/653,632  
;; FILING DATE: 24-MAY-1996  
;; ATTORNEY/AGENT INFORMATION:  
;; NAME: MCCORMACK, MYRA M.  
;; REGISTRATION NUMBER: 36,602  
;; REFERENCE/DOCKET NUMBER: 110.00330220  
;; TELECOMMUNICATION INFORMATION:  
;; TELEPHONE: 612-305-1225  
;; TELEFAX: 612-305-1228  
;; INFORMATION FOR SEQ ID NO: 13:  
;; SEQUENCE CHARACTERISTICS:  
;; LENGTH: 33 amino acids  
;; TYPE: amino acid  
;; STRANDEDNESS: single  
;; TOPOLOGY: linear  
;; MOLECULE TYPE: peptide  
;; SEQUENCE DESCRIPTION: SEQ ID NO: 13:  
US-10-300-083-13

Query Match 30.9%; Score 30; DB 14; Length 33;  
Best Local Similarity 60.0%; Pred. No. 9.7e+02;  
Matches 6; Conservative 1; Mismatches 3; Indels 0; Gaps 0;

QY 4 HLNSKIAFKI 13  
||:|:|:  
DB 15 HLKWKIIFKL 24

RESULT 66  
US-10-424-599-154283  
; Sequence 154283, Application US/10424599  
; Publication No. US20040031072A1  
; GENERAL INFORMATION:  
; APPLICANT: La Rosa Thomas J  
; APPLICANT: Kovalic David K  
; APPLICANT: Zhou Yihua  
; APPLICANT: Cao Yongwei  
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With  
; TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement  
; FILE REFERENCE: 38-21(53223)B  
; CURRENT APPLICATION NUMBER: US/10/424,599  
; CURRENT FILING DATE: 2003-04-28  
; NUMBER OF SEQ ID NOS: 285684  
; SEQ ID NO 154283  
; LENGTH: 34  
; TYPE: PRT  
; ORGANISM: Glycine max  
; FEATURE:  
; OTHER INFORMATION: Clone ID: PAT\_MRT3847\_110339C.1.pap  
US-10-424-599-154283

Query Match 30.9%; Score 30; DB 15; Length 34;  
Best Local Similarity 45.5%; Pred. No. 1e+03;  
Matches 5; Conservative 4; Mismatches 2; Indels 0; Gaps 0;

QY 3 NHLNSKIAFKI 13  
||:|:|:  
DB 13 NYLDSITWTFKL 23

```
RESULT 67
US-10-437-963-139274
; Sequence 139274, Application US/10437963
; Publication No. US20040123343A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa, Thomas J.
; APPLICANT: Kovalic, David K.
; APPLICANT: Zhou, Yihua
; APPLICANT: Cao, Yongwei
; APPLICANT: Wu, Wei
; APPLICANT: Boukharov, Andrey A.
; APPLICANT: Barbazuk, Brad
; APPLICANT: Li, Ping
; TITLE OF INVENTION: Rice Nucleic Acid Molecules and Other Molecules Associated With
; TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement
; FILE REFERENCE: 38-21(53221)B
; CURRENT APPLICATION NUMBER: US/10/437,963
; CURRENT FILING DATE: 2003-05-14
; NUMBER OF SEQ ID NOS: 204966
; SEQ ID NO 139274
; LENGTH: 36
; TYPE: PRT
; ORGANISM: Oryza sativa
; FEATURE:
; OTHER INFORMATION: Clone ID: PAT_MRT4530_40582C.1.pap
US-10-437-963-139274

Query Match          30.9%; Score 30; DB 16; Length 36;
Best Local Similarity 42.9%; Pred. No. 1.1e+03;
Matches 6; Conservative 2; Mismatches 6; Indels 0; Gaps 0;

QY      4 HLNSKIAPKIVSQE 17
DB      3 HLPQYILFKLMQDE 16

RESULT 68
US-10-437-963-176739
; Sequence 176739, Application US/10437963
; Publication No. US20040123343A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa, Thomas J.
; APPLICANT: Kovalic, David K.
; APPLICANT: Zhou, Yihua
; APPLICANT: Cao, Yongwei
; APPLICANT: Wu, Wei
; APPLICANT: Boukharov, Andrey A.
; APPLICANT: Barbazuk, Brad
; APPLICANT: Li, Ping
; TITLE OF INVENTION: Rice Nucleic Acid Molecules and Other Molecules Associated With
; TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement
; FILE REFERENCE: 38-21(53221)B
; CURRENT APPLICATION NUMBER: US/10/437,963
; CURRENT FILING DATE: 2003-05-14
; NUMBER OF SEQ ID NOS: 204966
; SEQ ID NO 176739
; LENGTH: 36
; TYPE: PRT
; ORGANISM: Oryza sativa
; FEATURE:
; OTHER INFORMATION: Clone ID: PAT_MRT4530_7445C.1.pap
US-10-437-963-176739

Query Match          30.9%; Score 30; DB 16; Length 36;
Best Local Similarity 35.7%; Pred. No. 1.1e+03;
Matches 5; Conservative 5; Mismatches 5; Indels 0; Gaps 0;

QY      3 NHLNSKIAPKIVSQ 16
DB      13 DHRSSQVAFSTYSE 26

RESULT 69
US-09-809-391-545
; Sequence 545, Application US/09809391
; Publication No. US20030049618A1
; GENERAL INFORMATION:
; APPLICANT: Ruben et al.
; TITLE OF INVENTION: 186 Human Secreted proteins
; FILE REFERENCE: PZ002P2
; CURRENT APPLICATION NUMBER: US/09/809,391
; CURRENT FILING DATE: 2001-03-16
; Prior application data removed - consult PALM or file wrapper
; NUMBER OF SEQ ID NOS: 761
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 545
; LENGTH: 39
; TYPE: PRT
; ORGANISM: Homo sapiens
US-09-809-391-545

Query Match          30.9%; Score 30; DB 10; Length 39;
Best Local Similarity 58.3%; Pred. No. 1.2e+03;
Matches 7; Conservative 1; Mismatches 4; Indels 0; Gaps 0;

QY      3 NHLNSKIAPKIV 14
DB      27 NHLAFRIILFFIV 38

RESULT 70
US-09-882-171-545
; Sequence 545, Application US/09882171
; Publication No. US20030175858A1
; GENERAL INFORMATION:
; APPLICANT: Ruben et al.
; TITLE OF INVENTION: 186 Human Secreted proteins
; FILE REFERENCE: PZ002P2
; CURRENT APPLICATION NUMBER: US/09/882,171
; CURRENT FILING DATE: 2001-06-18
; Prior application data removed - consult PALM or file wrapper
; NUMBER OF SEQ ID NOS: 761
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 545
; LENGTH: 39
; TYPE: PRT
; ORGANISM: Homo sapiens
US-09-882-171-545

Query Match          30.9%; Score 30; DB 10; Length 39;
Best Local Similarity 58.3%; Pred. No. 1.2e+03;
Matches 7; Conservative 1; Mismatches 4; Indels 0; Gaps 0;

QY      3 NHLNSKIAPKIV 14
DB      27 NHLAFRIILFFIV 38
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,	PRIOR FILING DATE:	1997-08-22
,	PRIOR APPLICATION NUMBER:	60/056,637
,	PRIOR FILING DATE:	1997-08-22
,	PRIOR APPLICATION NUMBER:	60/056,903
,	PRIOR FILING DATE:	1997-08-22
,	PRIOR APPLICATION NUMBER:	60/056,888
,	PRIOR FILING DATE:	1997-08-22
,	PRIOR APPLICATION NUMBER:	60/056,879
,	PRIOR FILING DATE:	1997-08-22
,	PRIOR APPLICATION NUMBER:	60/056,880
,	PRIOR FILING DATE:	1997-08-22
,	PRIOR APPLICATION NUMBER:	60/056,894
,	PRIOR FILING DATE:	1997-08-22
,	PRIOR APPLICATION NUMBER:	60/056,911
,	PRIOR FILING DATE:	1997-08-22
,	PRIOR APPLICATION NUMBER:	60/056,636
,	PRIOR FILING DATE:	1997-08-22
,	PRIOR APPLICATION NUMBER:	60/056,874
,	PRIOR FILING DATE:	1997-08-22
,	PRIOR APPLICATION NUMBER:	60/056,910
,	PRIOR FILING DATE:	1997-08-22
,	PRIOR APPLICATION NUMBER:	60/056,864
,	PRIOR FILING DATE:	1997-08-22
,	PRIOR APPLICATION NUMBER:	60/056,631
,	PRIOR FILING DATE:	1997-08-22
,	PRIOR APPLICATION NUMBER:	60/056,845
,	PRIOR FILING DATE:	1997-08-22
,	PRIOR APPLICATION NUMBER:	60/056,892
,	PRIOR FILING DATE:	1997-08-22
,	PRIOR APPLICATION NUMBER:	60/057,761
,	PRIOR FILING DATE:	1997-08-22
,	PRIOR APPLICATION NUMBER:	60/047,595
,	PRIOR FILING DATE:	1997-05-23
,	PRIOR APPLICATION NUMBER:	60/047,588
,	PRIOR FILING DATE:	1997-05-23
,	PRIOR APPLICATION NUMBER:	60/047,585
,	PRIOR FILING DATE:	1997-05-23
,	PRIOR APPLICATION NUMBER:	60/047,586
,	PRIOR FILING DATE:	1997-05-23
,	PRIOR APPLICATION NUMBER:	60/047,590
,	PRIOR FILING DATE:	1997-05-23
,	PRIOR APPLICATION NUMBER:	60/047,594
,	PRIOR FILING DATE:	1997-05-23
,	PRIOR APPLICATION NUMBER:	60/047,589
,	PRIOR FILING DATE:	1997-05-23
,	PRIOR APPLICATION NUMBER:	60/047,593
,	PRIOR FILING DATE:	1997-05-23
,	PRIOR APPLICATION NUMBER:	60/047,614
,	PRIOR FILING DATE:	1997-05-23
,	PRIOR APPLICATION NUMBER:	60/043,578
,	PRIOR FILING DATE:	1997-04-11
,	PRIOR APPLICATION NUMBER:	60/043,576
,	PRIOR FILING DATE:	1997-04-11
,	PRIOR APPLICATION NUMBER:	60/047,501
,	PRIOR FILING DATE:	1997-05-23
,	PRIOR APPLICATION NUMBER:	60/043,670
,	PRIOR FILING DATE:	1997-04-11
,	PRIOR APPLICATION NUMBER:	60/056,632
,	PRIOR FILING DATE:	1997-08-22
,	PRIOR APPLICATION NUMBER:	60/056,664
,	PRIOR FILING DATE:	1997-08-22
,	PRIOR APPLICATION NUMBER:	60/056,876
,	PRIOR FILING DATE:	1997-08-22
,	PRIOR APPLICATION NUMBER:	60/056,881
,	PRIOR FILING DATE:	1997-08-22
,	PRIOR APPLICATION NUMBER:	60/056,909
,	PRIOR FILING DATE:	1997-08-22
,	PRIOR APPLICATION NUMBER:	60/056,875
,	PRIOR FILING DATE:	1997-08-22
,	PRIOR APPLICATION NUMBER:	60/056,862
,	PRIOR FILING DATE:	1997-08-22

; PRIOR APPLICATION NUMBER: 60/056,887  
; PRIOR FILING DATE: 1997-08-22  
; PRIOR APPLICATION NUMBER: 60/056,908  
; PRIOR FILING DATE: 1997-08-22  
; PRIOR APPLICATION NUMBER: 60/048,964  
; PRIOR FILING DATE: 1997-06-06  
; PRIOR APPLICATION NUMBER: 60/057,650  
; PRIOR FILING DATE: 1997-09-05  
; PRIOR APPLICATION NUMBER: 60/056,884  
; PRIOR FILING DATE: 1997-08-22  
; PRIOR APPLICATION NUMBER: 60/057,669  
; PRIOR FILING DATE: 1997-09-05

Query Match 30.9%; Score 30; DB 10; Length 39;  
Best Local Similarity 58.3%; Pred. No. 1.2e+03;  
Matches 7; Conservative 1; Mismatches 4; Indels 0; Gaps 0;

QY 3 NNLNSKIAFKIV 14  
|||:|:|:  
DB 27 NHLAFILPFIV 38

RESULT 71  
US-09-962-756-202  
; Sequence 202, Application US/09962756  
; Publication No. US20030195147A1  
; GENERAL INFORMATION:  
; APPLICANT: PILLUTLA, RENUKA  
; APPLICANT: BRISSETTE, RENEE  
; APPLICANT: BLUME, ARTHUR J.  
; APPLICANT: SCHAEFFER, LAUGE  
; APPLICANT: BRANDT, JAKOB  
; APPLICANT: GOLDSTEIN, NEIL I.  
; APPLICANT: SPETZLER, JANE  
; APPLICANT: OSTERGAARD, SOREN  
; APPLICANT: HANSEN, PER HERTZ  
; TITLE OF INVENTION: INSULIN AND IGF-1 RECEPTOR AGONISTS AND ANTAGONISTS  
; FILE REFERENCE: 1878-4051US1  
; CURRENT APPLICATION NUMBER: US/09/962,756  
; CURRENT FILING DATE: 2001-09-24  
; PRIOR APPLICATION NUMBER: 09/538,038  
; PRIOR FILING DATE: 2000-03-29  
; PRIOR APPLICATION NUMBER: 09/146,127  
; PRIOR FILING DATE: 1998-09-02  
; NUMBER OF SEQ ID NOS: 2227  
; SOFTWARE: PatentIn Ver. 2.1  
; SEQ ID NO 202  
; LENGTH: 39  
; TYPE: PRT  
; ORGANISM: Artificial Sequence  
; FEATURE:  
; OTHER INFORMATION: Description of Artificial Sequence: Synthetic  
US-09-962-756-202

Query Match 30.9%; Score 30; DB 10; Length 39;  
Best Local Similarity 60.0%; Pred. No. 1.2e+03;  
Matches 6; Conservative 2; Mismatches 2; Indels 0; Gaps 0;

QY 10 AFKIVSQEPA 19  
|:|:|:|:  
DB 14 ASKVSEPPA 23

RESULT 72  
US-10-132-585-4  
; Sequence 4, Application US/10132585  
; Publication No. US2003005523A1  
; GENERAL INFORMATION:  
; APPLICANT: Kapeller-Libermann, Rosanna  
; TITLE OF INVENTION: 26030, A HUMAN RHO-GAP FAMILY MEMBER AND  
; TITLE OF INVENTION: USES THEREFOR  
; FILE REFERENCE: MPI01-101PIRM

; CURRENT APPLICATION NUMBER: US/10/132,585  
; CURRENT FILING DATE: 2002-04-25  
; PRIOR APPLICATION NUMBER: 60/286,581  
; PRIOR FILING DATE: 2001-04-25  
; NUMBER OF SEQ ID NOS: 6  
; SOFTWARE: FastSeq for Windows Version 4.0  
; SEQ ID NO 4  
; LENGTH: 39  
; TYPE: PRT  
; ORGANISM: unknown  
; FEATURE:  
; OTHER INFORMATION: PFAM consensus rhoGAP domain  
US-10-132-585-4

Query Match 30.9%; Score 30; DB 14; Length 39;  
Best Local Similarity 50.0%; Pred. No. 1.2e+03;  
Matches 5; Conservative 2; Mismatches 3; Indels 0; Gaps 0;

QY 4 HLNKIAFKI 13  
|||:|:|:  
DB 30 HLNMLAFPL 39

RESULT 73  
US-10-106-698-5006  
; Sequence 5006, Application US/10106698  
; Publication No. US20030109690A1  
; GENERAL INFORMATION:  
; APPLICANT: Ruben et al.  
; TITLE OF INVENTION: Colon and Colon Cancer Associated Polynucleotides and Polypeptides  
; FILE REFERENCE: PA005P1  
; CURRENT APPLICATION NUMBER: US/10/106,698  
; CURRENT FILING DATE: 2002-03-27  
; PRIOR APPLICATION NUMBER: PCT/US00/26524  
; PRIOR FILING DATE: 2000-09-28  
; PRIOR APPLICATION NUMBER: US 60/157,137  
; PRIOR FILING DATE: 1999-09-29  
; PRIOR APPLICATION NUMBER: US 60/163,280  
; PRIOR FILING DATE: 1999-11-03  
; NUMBER OF SEQ ID NOS: 8564  
; SOFTWARE: PatentIn Ver. 3.0  
; SEQ ID NO 5006  
; LENGTH: 39  
; TYPE: PRT  
; ORGANISM: Homo sapiens  
US-10-106-698-5006

Query Match 30.9%; Score 30; DB 14; Length 39;  
Best Local Similarity 42.9%; Pred. No. 1.2e+03;  
Matches 6; Conservative 4; Mismatches 4; Indels 0; Gaps 0;

QY 1 EPNHLSKIAFKIV 14  
|:|:|:|:|:  
DB 19 KPHYLNIKUPNNIV 32

RESULT 74  
US-10-164-861-545  
; Sequence 545, Application US/10164861  
; Publication No. US2003025248A1  
; GENERAL INFORMATION:  
; APPLICANT: Rosen et al.  
; TITLE OF INVENTION: 186 Human Secreted proteins  
; FILE REFERENCE: PZ002P1  
; CURRENT APPLICATION NUMBER: US/10/164,861  
; CURRENT FILING DATE: 2002-06-10  
; PRIOR APPLICATION NUMBER: US/09/149,476  
; PRIOR FILING DATE: 1998-09-08  
; PRIOR APPLICATION NUMBER: PCT/US98/04493  
; PRIOR FILING DATE: 1998-03-06  
; NUMBER OF SEQ ID NOS: 757  
; SOFTWARE: PatentIn Ver. 2.0  
; SEQ ID NO 545

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; LENGTH: 39
; TYPE: PRT
; ORGANISM: Homo sapiens
US-10-164-861-545

Query Match      30.9%; Score 30; DB 14; Length 39;
Best Local Similarity 58.3%; Pred. No. 1.2e+03;
Matches 7; Conservative 1; Mismatches 4; Indels 0; Gaps 0;

Qy      3 NHLNSKIAFKIV 14
      |||||:|:|
Db      27 NHLAFRLFFIV 38

RESULT 75
US-10-253-471-202
; Sequence 202, Application US/10253471
; Publication No. US20030236190A1
; GENERAL INFORMATION:
; APPLICANT: PILLUTLA, RENUKA et al.
; TITLE OF INVENTION: INSULIN AND IGF-1 RECEPTOR AGONISTS AND ANTAGONISTS
; FILE REFERENCE: 1878-4057
; CURRENT APPLICATION NUMBER: US/10/253,471
; CURRENT FILING DATE: 2002-09-24
; PRIOR APPLICATION NUMBER: 09/962,756
; PRIOR FILING DATE: 2001-09-24
; PRIOR APPLICATION NUMBER: 09/538,038
; PRIOR FILING DATE: 2000-03-29
; PRIOR APPLICATION NUMBER: 09/146,127
; PRIOR FILING DATE: 1998-09-02
; NUMBER OF SEQ ID NOS: 2227
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 202
; LENGTH: 39
; TYPE: PRT
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence: Synthetic
US-10-253-471-202

Query Match      30.9%; Score 30; DB 14; Length 39;
Best Local Similarity 60.0%; Pred. No. 1.2e+03;
Matches 6; Conservative 2; Mismatches 2; Indels 0; Gaps 0;

Qy      10 AFKIVSQEPA 19
      |||||:|:|
Db      14 ASKVSEPPA 23

RESULT 76
US-10-253-493-202
; Sequence 202, Application US/10253493
; Publication No. US20040023887A1
; GENERAL INFORMATION:
; APPLICANT: PILLUTLA, RENUKA et al.
; TITLE OF INVENTION: INSULIN AND IGF-1 RECEPTOR AGONISTS AND ANTAGONISTS
; FILE REFERENCE: 1878-4056
; CURRENT APPLICATION NUMBER: US/10/253,493
; CURRENT FILING DATE: 2002-09-24
; PRIOR APPLICATION NUMBER: 09/962,756
; PRIOR FILING DATE: 2001-09-24
; PRIOR APPLICATION NUMBER: 09/538,038
; PRIOR FILING DATE: 2000-03-29
; PRIOR APPLICATION NUMBER: 09/146,127
; PRIOR FILING DATE: 1998-09-02
; NUMBER OF SEQ ID NOS: 2227
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 202
; LENGTH: 39
; TYPE: PRT
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence: Synthetic
US-10-253-493-202

; OTHER INFORMATION: Description of Artificial Sequence: Synthetic
; OTHER INFORMATION: peptide
US-10-253-493-202

Query Match      30.9%; Score 30; DB 15; Length 39;
Best Local Similarity 60.0%; Pred. No. 1.2e+03;
Matches 6; Conservative 2; Mismatches 2; Indels 0; Gaps 0;

Qy      10 AFKIVSQEPA 19
      |||||:|:|
Db      14 ASKVSEPPA 23

RESULT 77
US-09-809-391-381
; Sequence 381, Application US/09809391
; Publication No. US20030049618A1
; GENERAL INFORMATION:
; APPLICANT: Ruben et al.
; TITLE OF INVENTION: 186 Human Secreted proteins
; FILE REFERENCE: P2002P2
; CURRENT APPLICATION NUMBER: US/09/809,391
; CURRENT FILING DATE: 2001-03-16
; Prior application data removed - consult PALM or file wrapper
; NUMBER OF SEQ ID NOS: 761
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 381
; LENGTH: 40
; TYPE: PRT
; ORGANISM: Homo sapiens
; FEATURE:
; NAME/KEY: SITE
; LOCATION: (40)
; OTHER INFORMATION: Xaa equals stop translation
US-09-809-391-381

Query Match      30.9%; Score 30; DB 10; Length 40;
Best Local Similarity 58.3%; Pred. No. 1.2e+03;
Matches 7; Conservative 1; Mismatches 4; Indels 0; Gaps 0;

Qy      3 NHLNSKIAFKIV 14
      |||||:|:|
Db      27 NHLAFRLFFIV 38

RESULT 78
US-09-882-171-381
; Sequence 381, Application US/09882171
; Publication No. US20030175858A1
; GENERAL INFORMATION:
; APPLICANT: Ruben et al.
; TITLE OF INVENTION: 186 Human Secreted proteins
; FILE REFERENCE: P2002P2
; CURRENT APPLICATION NUMBER: US/09/882,171
; CURRENT FILING DATE: 2001-06-18
; PRIOR APPLICATION NUMBER: 09/809,391
; PRIOR FILING DATE: 2001-03-16
; PRIOR APPLICATION NUMBER: 09/149,476
; PRIOR FILING DATE: 1998-09-08
; PRIOR APPLICATION NUMBER: PCT/US98/04493
; PRIOR FILING DATE: 1998-03-06
; PRIOR APPLICATION NUMBER: 60/040,162
; PRIOR FILING DATE: 1997-03-07
; PRIOR APPLICATION NUMBER: 60/040,333
; PRIOR FILING DATE: 1997-03-07
; PRIOR APPLICATION NUMBER: 60/038,621
; PRIOR FILING DATE: 1997-03-07
; PRIOR APPLICATION NUMBER: 60/040,626
; PRIOR FILING DATE: 1997-03-07
; PRIOR APPLICATION NUMBER: 60/040,334
; PRIOR FILING DATE: 1997-03-07
; PRIOR APPLICATION NUMBER: 60/040,336
; PRIOR FILING DATE: 1997-03-07
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1 PRIOR APPLICATION NUMBER: 60/040,163  
2 PRIOR FILING DATE: 1997-03-07  
3 PRIOR APPLICATION NUMBER: 60/047,600  
4 PRIOR FILING DATE: 1997-05-23  
5 PRIOR APPLICATION NUMBER: 60/047,615  
6 PRIOR FILING DATE: 1997-05-23  
7 PRIOR APPLICATION NUMBER: 60/047,597  
8 PRIOR FILING DATE: 1997-05-23  
9 PRIOR APPLICATION NUMBER: 60/047,502  
10 PRIOR FILING DATE: 1997-05-23  
11 PRIOR APPLICATION NUMBER: 60/047,633  
12 PRIOR FILING DATE: 1997-05-23  
13 PRIOR APPLICATION NUMBER: 60/047,583  
14 PRIOR FILING DATE: 1997-05-23  
15 PRIOR APPLICATION NUMBER: 60/047,617  
16 PRIOR FILING DATE: 1997-05-23  
17 PRIOR APPLICATION NUMBER: 60/047,618  
18 PRIOR FILING DATE: 1997-05-23  
19 PRIOR APPLICATION NUMBER: 60/047,503  
20 PRIOR FILING DATE: 1997-05-23  
21 PRIOR APPLICATION NUMBER: 60/047,592  
22 PRIOR FILING DATE: 1997-05-23  
23 PRIOR APPLICATION NUMBER: 60/047,581  
24 PRIOR FILING DATE: 1997-05-23  
25 PRIOR APPLICATION NUMBER: 60/047,584  
26 PRIOR FILING DATE: 1997-05-23  
27 PRIOR APPLICATION NUMBER: 60/047,500  
28 PRIOR FILING DATE: 1997-05-23  
29 PRIOR APPLICATION NUMBER: 60/047,587  
30 PRIOR FILING DATE: 1997-05-23  
31 PRIOR APPLICATION NUMBER: 60/047,492  
32 PRIOR FILING DATE: 1997-05-23  
33 PRIOR APPLICATION NUMBER: 60/047,598  
34 PRIOR FILING DATE: 1997-05-23  
35 PRIOR APPLICATION NUMBER: 60/047,613  
36 PRIOR FILING DATE: 1997-05-23  
37 PRIOR APPLICATION NUMBER: 60/047,582  
38 PRIOR FILING DATE: 1997-05-23  
39 PRIOR APPLICATION NUMBER: 60/047,596  
40 PRIOR FILING DATE: 1997-05-23  
41 PRIOR APPLICATION NUMBER: 60/047,612  
42 PRIOR FILING DATE: 1997-05-23  
43 PRIOR APPLICATION NUMBER: 60/047,632  
44 PRIOR FILING DATE: 1997-05-23  
45 PRIOR APPLICATION NUMBER: 60/047,601  
46 PRIOR FILING DATE: 1997-05-23  
47 PRIOR APPLICATION NUMBER: 60/043,580  
48 PRIOR FILING DATE: 1997-04-11  
49 PRIOR APPLICATION NUMBER: 60/043,568  
50 PRIOR FILING DATE: 1997-04-11  
51 PRIOR APPLICATION NUMBER: 60/043,314  
52 PRIOR FILING DATE: 1997-04-11  
53 PRIOR APPLICATION NUMBER: 60/043,569  
54 PRIOR FILING DATE: 1997-04-11  
55 PRIOR APPLICATION NUMBER: 60/043,311  
56 PRIOR FILING DATE: 1997-04-11  
57 PRIOR APPLICATION NUMBER: 60/043,671  
58 PRIOR FILING DATE: 1997-04-11  
59 PRIOR APPLICATION NUMBER: 60/043,674  
60 PRIOR FILING DATE: 1997-04-11  
61 PRIOR APPLICATION NUMBER: 60/043,669  
62 PRIOR FILING DATE: 1997-04-11  
63 PRIOR APPLICATION NUMBER: 60/043,312  
64 PRIOR FILING DATE: 1997-04-11  
65 PRIOR APPLICATION NUMBER: 60/043,313  
66 PRIOR FILING DATE: 1997-04-11  
67 PRIOR APPLICATION NUMBER: 60/043,672  
68 PRIOR FILING DATE: 1997-04-11  
69 PRIOR APPLICATION NUMBER: 60/043,315  
70 PRIOR FILING DATE: 1997-04-11  
71 PRIOR APPLICATION NUMBER: 60/048,974  
72 PRIOR FILING DATE: 1997-06-06  
73 PRIOR APPLICATION NUMBER: 60/056,886

74 PRIOR FILING DATE: 1997-08-22  
75 PRIOR APPLICATION NUMBER: 60/056,877  
76 PRIOR FILING DATE: 1997-08-22  
77 PRIOR APPLICATION NUMBER: 60/056,889  
78 PRIOR FILING DATE: 1997-08-22  
79 PRIOR APPLICATION NUMBER: 60/056,893  
80 PRIOR FILING DATE: 1997-08-22  
81 PRIOR APPLICATION NUMBER: 60/056,630  
82 PRIOR FILING DATE: 1997-08-22  
83 PRIOR APPLICATION NUMBER: 60/056,878  
84 PRIOR FILING DATE: 1997-08-22  
85 PRIOR APPLICATION NUMBER: 60/056,662  
86 PRIOR FILING DATE: 1997-08-22  
87 PRIOR APPLICATION NUMBER: 60/056,872  
88 PRIOR FILING DATE: 1997-08-22  
89 PRIOR APPLICATION NUMBER: 60/056,882  
90 PRIOR FILING DATE: 1997-08-22  
91 PRIOR APPLICATION NUMBER: 60/056,637  
92 PRIOR FILING DATE: 1997-08-22  
93 PRIOR APPLICATION NUMBER: 60/056,903  
94 PRIOR FILING DATE: 1997-08-22  
95 PRIOR APPLICATION NUMBER: 60/056,888  
96 PRIOR FILING DATE: 1997-08-22  
97 PRIOR APPLICATION NUMBER: 60/056,879  
98 PRIOR FILING DATE: 1997-08-22  
99 PRIOR APPLICATION NUMBER: 60/056,880  
100 PRIOR FILING DATE: 1997-08-22  
101 PRIOR APPLICATION NUMBER: 60/056,894  
102 PRIOR FILING DATE: 1997-08-22  
103 PRIOR APPLICATION NUMBER: 60/056,911  
104 PRIOR FILING DATE: 1997-08-22  
105 PRIOR APPLICATION NUMBER: 60/056,636  
106 PRIOR FILING DATE: 1997-08-22  
107 PRIOR APPLICATION NUMBER: 60/056,874  
108 PRIOR FILING DATE: 1997-08-22  
109 PRIOR APPLICATION NUMBER: 60/056,910  
110 PRIOR FILING DATE: 1997-08-22  
111 PRIOR APPLICATION NUMBER: 60/056,864  
112 PRIOR FILING DATE: 1997-08-22  
113 PRIOR APPLICATION NUMBER: 60/056,631  
114 PRIOR FILING DATE: 1997-08-22  
115 PRIOR APPLICATION NUMBER: 60/056,845  
116 PRIOR FILING DATE: 1997-08-22  
117 PRIOR APPLICATION NUMBER: 60/056,892  
118 PRIOR FILING DATE: 1997-08-22  
119 PRIOR APPLICATION NUMBER: 60/057,761  
120 PRIOR FILING DATE: 1997-08-22  
121 PRIOR APPLICATION NUMBER: 60/047,595  
122 PRIOR FILING DATE: 1997-05-23  
123 PRIOR APPLICATION NUMBER: 60/047,599  
124 PRIOR FILING DATE: 1997-05-23  
125 PRIOR APPLICATION NUMBER: 60/047,588  
126 PRIOR FILING DATE: 1997-05-23  
127 PRIOR APPLICATION NUMBER: 60/047,585  
128 PRIOR FILING DATE: 1997-05-23  
129 PRIOR APPLICATION NUMBER: 60/047,586  
130 PRIOR FILING DATE: 1997-05-23  
131 PRIOR APPLICATION NUMBER: 60/047,590  
132 PRIOR FILING DATE: 1997-05-23  
133 PRIOR APPLICATION NUMBER: 60/047,594  
134 PRIOR FILING DATE: 1997-05-23  
135 PRIOR APPLICATION NUMBER: 60/047,589  
136 PRIOR FILING DATE: 1997-05-23  
137 PRIOR APPLICATION NUMBER: 60/047,593  
138 PRIOR FILING DATE: 1997-05-23  
139 PRIOR APPLICATION NUMBER: 60/047,614  
140 PRIOR FILING DATE: 1997-05-23  
141 PRIOR APPLICATION NUMBER: 60/043,578  
142 PRIOR FILING DATE: 1997-04-11  
143 PRIOR APPLICATION NUMBER: 60/043,576  
144 PRIOR FILING DATE: 1997-04-11  
145 PRIOR APPLICATION NUMBER: 60/047,501  
146 PRIOR FILING DATE: 1997-05-23



```
; PRIOR APPLICATION NUMBER: 60/043,670
; PRIOR FILING DATE: 1997-04-11
; PRIOR APPLICATION NUMBER: 60/056,632
; PRIOR FILING DATE: 1997-08-22
; PRIOR APPLICATION NUMBER: 60/056,664
; PRIOR FILING DATE: 1997-08-22
; PRIOR APPLICATION NUMBER: 60/056,876
; PRIOR FILING DATE: 1997-08-22
; PRIOR APPLICATION NUMBER: 60/056,881
; PRIOR FILING DATE: 1997-08-22
; PRIOR APPLICATION NUMBER: 60/056,909
; PRIOR FILING DATE: 1997-08-22
; PRIOR APPLICATION NUMBER: 60/056,875
; PRIOR FILING DATE: 1997-08-22
; PRIOR APPLICATION NUMBER: 60/056,862
; PRIOR FILING DATE: 1997-08-22
; PRIOR APPLICATION NUMBER: 60/056,887
; PRIOR FILING DATE: 1997-08-22
; PRIOR APPLICATION NUMBER: 60/056,908
; PRIOR FILING DATE: 1997-08-22
; PRIOR APPLICATION NUMBER: 60/048,964
; PRIOR FILING DATE: 1997-06-06
; PRIOR APPLICATION NUMBER: 60/057,650
; PRIOR FILING DATE: 1997-09-05
; PRIOR APPLICATION NUMBER: 60/056,884
; PRIOR FILING DATE: 1997-08-22
; PRIOR APPLICATION NUMBER: 60/057,669
; PRIOR FILING DATE: 1997-09-05
```

```
Query Match          30.9%; Score 30; DB 10; Length 40;
Best Local Similarity 58.3%; Pred. No. 1.2e+03;
Matches 7; Conservative 1; Mismatches 4; Indels 0; Gaps 0;
```

```
Qy 3 NHLNSKIAFKIV 14
    ||| : |||
Db 27 NHLAFRIILFFIV 38
```

## RESULT 79

```
US-10-144-259-29
; Sequence 29, Application US/10144259
; Publication No. US20030109691A1
; GENERAL INFORMATION:
; APPLICANT: Arnaout, M. Amin
; APPLICANT: Li, Rui
; APPLICANT: Xiong, Jian-Ping
; TITLE OF INVENTION: VARIANT INTEGRIN POLYPEPTIDES AND USES THEREOF
; FILE REFERENCE: 00786-548001
; CURRENT APPLICATION NUMBER: US/10/144,259
; CURRENT FILING DATE: 2002-09-04
; PRIOR APPLICATION NUMBER: US 09/758,493
; PRIOR FILING DATE: 2001-01-11
; PRIOR APPLICATION NUMBER: US 60/221,950
; PRIOR FILING DATE: 2000-07-31
; NUMBER OF SEQ ID NOS: 30
; SOFTWARE: FastSEQ for Windows Version 4.0
; SEQ ID NO 29
; LENGTH: 40
; TYPE: PRT
; ORGANISM: Homo sapiens
US-10-144-259-29
```

```
Query Match          30.9%; Score 30; DB 14; Length 40;
Best Local Similarity 46.2%; Pred. No. 1.2e+03;
Matches 6; Conservative 3; Mismatches 4; Indels 0; Gaps 0;
```

```
Qy 5 LNSKIAFKIVSQE 17
    ||| : |||
Db 7 LLSKLYNIISME 19
```

## RESULT 80

```
US-10-164-861-381
```

```
; Sequence 381, Application US/10164861
; Publication No. US20030225248A1
; GENERAL INFORMATION:
; APPLICANT: Rosen et al.
; TITLE OF INVENTION: 186 Human Secreted proteins
; FILE REFERENCE: PZ002P1
; CURRENT APPLICATION NUMBER: US/10/164,861
; CURRENT FILING DATE: 2002-06-10
; PRIOR APPLICATION NUMBER: US/09/149,476
; PRIOR FILING DATE: 1998-09-08
; PRIOR APPLICATION NUMBER: PCT/US98/04493
; PRIOR FILING DATE: 1998-03-06
; NUMBER OF SEQ ID NOS: 757
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 381
; LENGTH: 40
; TYPE: PRT
; ORGANISM: Homo sapiens
; FEATURE:
; NAME/KEY: SITE
; LOCATION: (40)
; OTHER INFORMATION: Xaa equals stop translation
US-10-164-861-381
```

```
Query Match          30.9%; Score 30; DB 14; Length 40;
Best Local Similarity 58.3%; Pred. No. 1.2e+03;
Matches 7; Conservative 1; Mismatches 4; Indels 0; Gaps 0;
```

```
Qy 3 NHLNSKIAFKIV 14
    ||| : |||
Db 27 NHLAFRIILFFIV 38
```

## RESULT 81

```
US-10-424-599-267336
; Sequence 267336, Application US/10424599
; Publication No. US20040031072A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa Thomas J
; APPLICANT: Kovalic David K
; APPLICANT: Zhou Yihua
; APPLICANT: Cao Yongwei
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated with
; FILE REFERENCE: 38-21(53223)B
; CURRENT APPLICATION NUMBER: US/10/424,599
; CURRENT FILING DATE: 2003-04-28
; NUMBER OF SEQ ID NOS: 285684
; SEQ ID NO 267336
; LENGTH: 40
; TYPE: PRT
; ORGANISM: Glycine max
; FEATURE:
; OTHER INFORMATION: Clone ID: PAT_MRT3847_83426C.1.pep
US-10-424-599-267336
```

```
Query Match          30.9%; Score 30; DB 15; Length 40;
Best Local Similarity 85.7%; Pred. No. 1.2e+03;
Matches 6; Conservative 1; Mismatches 0; Indels 0; Gaps 0;
```

```
Qy 9 IAFKIVS 15
    : |||
Db 9 VAFKIVS 15
```

## RESULT 82

```
US-10-424-599-278809
; Sequence 278809, Application US/10424599
; Publication No. US20040031072A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa Thomas J
; APPLICANT: Kovalic David K
; APPLICANT: Zhou Yihua
```

; APPLICANT: Cao Yongwei  
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated with  
; FILE OF INVENTION: Plants and Uses Thereof for Plant Improvement  
; FILE REFERENCE: 38-21(53223)B  
; CURRENT APPLICATION NUMBER: US/10/424,599  
; CURRENT FILING DATE: 2003-04-28  
; NUMBER OF SEQ ID NOS: 285684  
; SEQ ID NO 278809  
; LENGTH: 44  
; TYPE: PRT  
; ORGANISM: Glycine max  
; FEATURE:  
; OTHER INFORMATION: Clone ID: PAT\_MRT3847\_93788C.1.pap  
US-10-424-599-278809

Query Match 30.9%; Score 30; DB 15; Length 44;  
Best Local Similarity 50.0%; Pred. No. 1.3e+03;  
Matches 6; Conservative 2; Mismatches 4; Indels 0; Gaps 0;

QY 3 NNLNSKIAFKIV 14  
||| ||| : :  
Db 10 NNLWSKSTWRVV 21

RESULT 83  
US-10-437-963-154105  
; Sequence 154105, Application US/10437963  
; Publication No. US20040123343A1  
; GENERAL INFORMATION:  
; APPLICANT: La Rosa, Thomas J.  
; APPLICANT: Kovalic, David K.  
; APPLICANT: Zhou, Yihua  
; APPLICANT: Cao, Yongwei  
; APPLICANT: Wu, Wei  
; APPLICANT: Boukharov, Andrey A.  
; APPLICANT: Barbazuk, Brad  
; APPLICANT: Li, Ping  
; TITLE OF INVENTION: Rice Nucleic Acid Molecules and Other Molecules Associated with  
; FILE OF INVENTION: Plants and Uses Thereof for Plant Improvement  
; FILE REFERENCE: 38-21(53221)B  
; CURRENT APPLICATION NUMBER: US/10/437,963  
; CURRENT FILING DATE: 2003-05-14  
; NUMBER OF SEQ ID NOS: 204966  
; SEQ ID NO 154105  
; LENGTH: 44  
; TYPE: PRT  
; ORGANISM: Oryza sativa  
; FEATURE:  
; OTHER INFORMATION: Clone ID: PAT\_MRT4530\_53998C.1.pap  
US-10-437-963-154105

Query Match 30.9%; Score 30; DB 16; Length 44;  
Best Local Similarity 45.5%; Pred. No. 1.3e+03;  
Matches 5; Conservative 2; Mismatches 4; Indels 0; Gaps 0;

QY 2 PNLNSKIAFK 12  
||| | : :  
Db 2 PNLFSVKTYK 12

RESULT 84  
US-10-424-599-228808  
; Sequence 228808, Application US/10424599  
; Publication No. US20040031072A1  
; GENERAL INFORMATION:  
; APPLICANT: La Rosa Thomas J  
; APPLICANT: Kovalic David K  
; APPLICANT: Zhou Yihua  
; APPLICANT: Cao Yongwei  
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated with  
; FILE OF INVENTION: Plants and Uses Thereof for Plant Improvement  
; FILE REFERENCE: 38-21(53223)B  
; CURRENT APPLICATION NUMBER: US/10/424,599

; CURRENT FILING DATE: 2003-04-28  
; NUMBER OF SEQ ID NOS: 285684  
; SEQ ID NO 228808  
; LENGTH: 45  
; TYPE: PRT  
; ORGANISM: Glycine max  
; FEATURE:  
; NAME/KEY: unsure  
; LOCATION: (1)..(45)  
; OTHER INFORMATION: unsure at all Xaa locations  
; FEATURE:  
; OTHER INFORMATION: Clone ID: PAT\_MRT3847\_48640C.1.pap  
US-10-424-599-228808

Query Match 30.9%; Score 30; DB 15; Length 45;  
Best Local Similarity 58.3%; Pred. No. 1.4e+03;  
Matches 7; Conservative 2; Mismatches 1; Indels 2; Gaps 1;

QY 3 NNLNSKIAFKIV 14  
||| : : :  
Db 7 NNL--TKLTFKIV 16

RESULT 85  
US-09-764-877-1428  
; Sequence 1428, Application US/09764877  
; Patent No. US20020147140A1  
; GENERAL INFORMATION:  
; APPLICANT: Rosen et al.  
; TITLE OF INVENTION: Nucleic Acids, Proteins, and Antibodies  
; FILE REFERENCE: PC005  
; CURRENT APPLICATION NUMBER: US/09/764,877  
; CURRENT FILING DATE: 2001-01-17  
; Prior application data removed - refer to PALM or file wrapper  
; NUMBER OF SEQ ID NOS: 4031  
; SOFTWARE: PatentIn Ver. 2.0  
; SEQ ID NO 1428  
; LENGTH: 46  
; TYPE: PRT  
; ORGANISM: Homo sapiens  
US-09-764-877-1428

Query Match 30.9%; Score 30; DB 9; Length 46;  
Best Local Similarity 41.7%; Pred. No. 1.4e+03;  
Matches 5; Conservative 5; Mismatches 2; Indels 0; Gaps 0;

QY 5 LNSKIAFKIVSQ 16  
| : : : : :  
Db 28 LSQEVAFKLSTQ 39

RESULT 86  
US-10-242-515-1428  
; Sequence 1428, Application US/10242515  
; Publication No. US20040009488A1  
; GENERAL INFORMATION:  
; APPLICANT: Rosen et al.  
; TITLE OF INVENTION: Nucleic Acids, Proteins, and Antibodies  
; FILE REFERENCE: PC005C1  
; CURRENT APPLICATION NUMBER: US/10/242,515  
; CURRENT FILING DATE: 2002-09-13  
; PRIOR APPLICATION NUMBER: 09/764,877  
; PRIOR FILING DATE: 2001-01-17  
; PRIOR APPLICATION NUMBER: 60/179,065  
; PRIOR FILING DATE: 2000-01-31  
; PRIOR APPLICATION NUMBER: 60/180,628  
; PRIOR FILING DATE: 2000-02-04  
; PRIOR APPLICATION NUMBER: 60/214,896  
; PRIOR FILING DATE: 2000-06-28  
; PRIOR APPLICATION NUMBER: 60/217,487  
; PRIOR FILING DATE: 2000-07-11  
; PRIOR APPLICATION NUMBER: 60/225,758  
; PRIOR FILING DATE: 2000-08-14

```
; PRIOR APPLICATION NUMBER: 60/220,963
; PRIOR FILING DATE: 2000-07-26
; PRIOR APPLICATION NUMBER: 60/217,496
; PRIOR FILING DATE: 2000-07-11
; PRIOR APPLICATION NUMBER: 60/225,447
; PRIOR FILING DATE: 2000-08-14
; PRIOR APPLICATION NUMBER: 60/218,290
; PRIOR FILING DATE: 2000-07-14
; Remaining Prior Application data removed - See File Wrapper or PALM.
; NUMBER OF SEQ ID NOS: 4031
; SOFTWARE: Patentin Ver. 2.0
; SEQ ID NO 1428
; LENGTH: 46
; TYPE: PRT
; ORGANISM: Homo sapiens
US-10-242-515-1428

Query Match          30.9%; Score 30; DB 15; Length 46;
Best Local Similarity 41.7%; Pred. No. 1.4e+03;
Matches 5; Conservative 5; Mismatches 2; Indels 0; Gaps 0;

Qy 5 LNSKIAFKIVSQ 16
Db 28 LSQEVAFKLSQ 39
|: ::|||: |:|

RESULT 87
US-10-424-599-264089
; Sequence 264089, Application US/10424599
; Publication No. US20040031072A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa Thomas J
; APPLICANT: Kovalic David K
; APPLICANT: Zhou Yihua
; APPLICANT: Cao Yongwei
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With
; FILE REFERENCE: 38-21(53223)B
; CURRENT APPLICATION NUMBER: US/10/424,599
; CURRENT FILING DATE: 2003-04-28
; NUMBER OF SEQ ID NOS: 285684
; SEQ ID NO 264089
; LENGTH: 47
; TYPE: PRT
; ORGANISM: Glycine max
; FEATURE:
; OTHER INFORMATION: Clone ID: PAT_MRT3847_80493C.1.pep
US-10-424-599-264089

Query Match          30.9%; Score 30; DB 15; Length 47;
Best Local Similarity 54.5%; Pred. No. 1.4e+03;
Matches 6; Conservative 1; Mismatches 4; Indels 0; Gaps 0;

Qy 3 NHLNSKIAFKI 13
Db 7 NKLEGLHFVKV 17
|: |||: |:|

RESULT 88
US-10-424-599-208160
; Sequence 208160, Application US/10424599
; Publication No. US20040031072A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa Thomas J
; APPLICANT: Kovalic David K
; APPLICANT: Zhou Yihua
; APPLICANT: Cao Yongwei
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With
; FILE REFERENCE: 38-21(53223)B
; CURRENT APPLICATION NUMBER: US/10/424,599
; CURRENT FILING DATE: 2003-04-28
; NUMBER OF SEQ ID NOS: 285684
```

```
; SEQ ID NO 208160
; LENGTH: 48
; TYPE: PRT
; ORGANISM: Glycine max
; FEATURE:
; OTHER INFORMATION: Clone ID: PAT_MRT3847_29995C.1.pep
US-10-424-599-208160

Query Match          30.9%; Score 30; DB 15; Length 48;
Best Local Similarity 42.9%; Pred. No. 1.5e+03;
Matches 6; Conservative 3; Mismatches 5; Indels 0; Gaps 0;

Qy 3 NHLNSKIAFKIVSQ 16
Db 33 NHPNSKNKFLVQNE 46
|: |||: |:|

RESULT 89
US-10-424-599-264585
; Sequence 264585, Application US/10424599
; Publication No. US20040031072A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa Thomas J
; APPLICANT: Kovalic David K
; APPLICANT: Zhou Yihua
; APPLICANT: Cao Yongwei
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With
; FILE REFERENCE: 38-21(53223)B
; CURRENT APPLICATION NUMBER: US/10/424,599
; CURRENT FILING DATE: 2003-04-28
; NUMBER OF SEQ ID NOS: 285684
; SEQ ID NO 264585
; LENGTH: 48
; TYPE: PRT
; ORGANISM: Glycine max
; FEATURE:
; OTHER INFORMATION: Clone ID: PAT_MRT3847_80940C.1.pep
US-10-424-599-264585

Query Match          30.9%; Score 30; DB 15; Length 48;
Best Local Similarity 54.5%; Pred. No. 1.5e+03;
Matches 6; Conservative 2; Mismatches 3; Indels 0; Gaps 0;

Qy 4 HLNSKIAFKIV 14
Db 26 HVISKIKLVV 36
|: |||: |:|

RESULT 90
US-10-424-599-231764
; Sequence 231764, Application US/10424599
; Publication No. US20040031072A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa Thomas J
; APPLICANT: Kovalic David K
; APPLICANT: Zhou Yihua
; APPLICANT: Cao Yongwei
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With
; FILE REFERENCE: 38-21(53223)B
; CURRENT APPLICATION NUMBER: US/10/424,599
; CURRENT FILING DATE: 2003-04-28
; NUMBER OF SEQ ID NOS: 285684
; SEQ ID NO 231764
; LENGTH: 49
; TYPE: PRT
; ORGANISM: Glycine max
; FEATURE:
; OTHER INFORMATION: Clone ID: PAT_MRT3847_51305C.1.pep
US-10-424-599-231764

Query Match          30.9%; Score 30; DB 15; Length 49;
```

Best Local Similarity 38.9%; Pred. No. 1.5e+03;  
Matches 7; Conservative 3; Mismatches 6; Indels 2; Gaps 1;

QY 3 NH--LNSKIAPKIVSQEP 18  
|||:::|  
Db 16 NHPFINTNSNFKVILLHP 33

## RESULT 91

US-10-424-599-239689  
; Sequence 239689, Application US/10424599  
; Publication No. US20040031072A1  
; GENERAL INFORMATION:  
; APPLICANT: La Rosa Thomas J  
; APPLICANT: Kovalic David K  
; APPLICANT: Zhou Yihua  
; APPLICANT: Cao Yongwei  
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With  
; TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement  
; FILE REFERENCE: 38-21(53223)B  
; CURRENT APPLICATION NUMBER: US/10/424,599  
; CURRENT FILING DATE: 2003-04-28  
; NUMBER OF SEQ ID NOS: 285684  
; SEQ ID NO 239689  
; LENGTH: 49  
; TYPE: PRT  
; ORGANISM: Glycine max  
; FEATURE:  
; OTHER INFORMATION: Clone ID: PAT\_MRT3847\_59464C.1.pap  
US-10-424-599-239689

Query Match 30.9%; Score 30; DB 15; Length 49;  
Best Local Similarity 83.3%; Pred. No. 1.5e+03;  
Matches 5; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 3 NHLNSK 8  
|||||  
Db 30 NHLNTX 35

## RESULT 92

US-10-424-599-244508  
; Sequence 244508, Application US/10424599  
; Publication No. US20040031072A1  
; GENERAL INFORMATION:  
; APPLICANT: La Rosa Thomas J  
; APPLICANT: Kovalic David K  
; APPLICANT: Zhou Yihua  
; APPLICANT: Cao Yongwei  
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With  
; TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement  
; FILE REFERENCE: 38-21(53223)B  
; CURRENT APPLICATION NUMBER: US/10/424,599  
; CURRENT FILING DATE: 2003-04-28  
; NUMBER OF SEQ ID NOS: 285684  
; SEQ ID NO 244508  
; LENGTH: 49  
; TYPE: PRT  
; ORGANISM: Glycine max  
; FEATURE:  
; OTHER INFORMATION: Clone ID: PAT\_MRT3847\_62818C.1.pap  
US-10-424-599-244508

Query Match 30.9%; Score 30; DB 15; Length 49;  
Best Local Similarity 46.7%; Pred. No. 1.5e+03;  
Matches 7; Conservative 2; Mismatches 6; Indels 0; Gaps 0;

QY 3 NHLNSKIAPKIVSQE 17  
|||:::|  
Db 22 NHEYKAPFPKILTOE 36

## RESULT 93

US-10-437-963-134140  
; Sequence 134140, Application US/10437963  
; Publication No. US20040123343A1  
; GENERAL INFORMATION:  
; APPLICANT: La Rosa, Thomas J.  
; APPLICANT: Kovalic, David K.  
; APPLICANT: Zhou, Yihua  
; APPLICANT: Cao, Yongwei  
; APPLICANT: Wu, Wei  
; APPLICANT: Boukharov, Andrey A.  
; APPLICANT: Boukharov, Brad  
; APPLICANT: Li, Ping  
; TITLE OF INVENTION: Rice Nucleic Acid Molecules and Other Molecules Associated With  
; TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement  
; FILE REFERENCE: 38-21(53221)B  
; CURRENT APPLICATION NUMBER: US/10/437,963  
; CURRENT FILING DATE: 2003-05-14  
; NUMBER OF SEQ ID NOS: 204966  
; SEQ ID NO 134140  
; LENGTH: 49  
; TYPE: PRT  
; ORGANISM: Oryza sativa  
; FEATURE:  
; OTHER INFORMATION: Clone ID: PAT\_MRT4530\_35942C.1.pap  
US-10-437-963-134140

Query Match 30.9%; Score 30; DB 16; Length 49;  
Best Local Similarity 50.0%; Pred. No. 1.5e+03;  
Matches 8; Conservative 1; Mismatches 3; Indels 4; Gaps 1;

QY 2 PNHLNSK---IAFKI 13  
|||||  
Db 33 PNHLIKKKRLSFAFKV 48

## RESULT 94

US-09-864-408A-4082  
; Sequence 4082, Application US/09864408A  
; Publication No. US20040009474A1  
; GENERAL INFORMATION:  
; APPLICANT: Leach, Martin D.  
; APPLICANT: Shimkets, Richard A.  
; TITLE OF INVENTION: No. US20040009474A1 Human Polynucleotides and Polypeptides Enc  
; FILE REFERENCE: 21402-012  
; CURRENT APPLICATION NUMBER: US/09/864,408A  
; CURRENT FILING DATE: 2001-05-24  
; PRIOR APPLICATION NUMBER: 60/206,690  
; PRIOR FILING DATE: 2000-05-24  
; NUMBER OF SEQ ID NOS: 9068  
; SOFTWARE: FastSeq for Windows Version 4.0  
; SEQ ID NO 4082  
; LENGTH: 50  
; TYPE: PRT  
; ORGANISM: Homo sapiens  
; FEATURE:  
; NAME/KEY: misc feature  
; LOCATION: (16)--(16)  
; OTHER INFORMATION: Wherein Xaa may be any naturally occurring amino acid  
US-09-864-408A-4082

Query Match 30.9%; Score 30; DB 11; Length 50;  
Best Local Similarity 37.5%; Pred. No. 1.5e+03;  
Matches 6; Conservative 3; Mismatches 7; Indels 0; Gaps 0;

QY 1 EPNHLNSKIAPKIVSQ 16  
|:::|  
Db 33 EEGMNGLIATLCGR 48

## RESULT 95

US-10-437-963-125237  
; Sequence 125237, Application US/10437963  
; Publication No. US20040123343A1

```
; GENERAL INFORMATION:
; APPLICANT: La Rosa, Thomas J.
; APPLICANT: Kovalic, David K.
; APPLICANT: Zhou, Yihua
; APPLICANT: Cao, Yongwei
; APPLICANT: Wu, Wei
; APPLICANT: Boukharov, Andrey A.
; APPLICANT: Barbazuk, Brad
; APPLICANT: Li, Ping
; TITLE OF INVENTION: Rice Nucleic Acid Molecules and Other Molecules Associated With
; TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement
; FILE REFERENCE: 38-21(53221)B
; CURRENT APPLICATION NUMBER: US/10/437.963
; CURRENT FILING DATE: 2003-05-14
; NUMBER OF SEQ ID NOS: 204966
; SEQ ID NO 125237
; LENGTH: 50
; TYPE: PRT
; ORGANISM: Oryza sativa
; FEATURE:
; OTHER INFORMATION: Clone ID: PAT_MRT4530_27900C.1.pep
US-10-437-963-125237

Query Match      30.9%; Score 30; DB 16; Length 50;
Best Local Similarity 21.4%; Pred. No. 1.5e+03;
Matches 3; Conservative 9; Mismatches 2; Indels 0; Gaps 0;

Qy      3 NHLNSKIAFKIVSQ 16
Db      20 SHLNRLSIQTIK 33

RESULT 96
US-10-164-359-12
; Sequence 12, Application US/10164359
; Publication No. US20030012776A1
; GENERAL INFORMATION:
; APPLICANT: Chin, Khew-Voon
; TITLE OF INVENTION: Nucleic Acid and Protein Expressed Thereby and Their Involvement
; TITLE OF INVENTION: Stress
; FILE REFERENCE: 601-1-108US
; CURRENT APPLICATION NUMBER: US/10/164,359
; CURRENT FILING DATE: 2002-08-06
; PRIOR FILING DATE: PCT/US00/33438
; PRIOR FILING DATE: 2000-12-07
; PRIOR FILING DATE: 60/169,418
; PRIOR FILING DATE: 1999-12-07
; NUMBER OF SEQ ID NOS: 18
; SOFTWARE: PatentIn version 3.1
; SEQ ID NO 12
; LENGTH: 24
; TYPE: PRT
; ORGANISM: Homo sapiens
US-10-164-359-12

Query Match      29.9%; Score 29; DB 14; Length 24;
Best Local Similarity 62.5%; Pred. No. 9.9e+02;
Matches 5; Conservative 1; Mismatches 2; Indels 0; Gaps 0;

Qy      2 PNHLNSKI 9
Db      11 PDHLNGHI 18

RESULT 97
US-10-413-785-6
; Sequence 6, Application US/10413785
; Publication No. US20030229906A1
; GENERAL INFORMATION:
; APPLICANT: Gelman et al.
; TITLE OF INVENTION: METHODS AND COMPOSITIONS FOR THE TREATMENT OF DISORDERS OF HIV
; TITLE OF INVENTION: INFECTION
; FILE REFERENCE: 29636/38269A
```

```
; CURRENT APPLICATION NUMBER: US/10/413,785
; CURRENT FILING DATE: 2003-04-14
; PRIOR APPLICATION NUMBER: US 60/372,557
; PRIOR FILING DATE: 2002-04-15
; NUMBER OF SEQ ID NOS: 15
; SOFTWARE: PatentIn version 3.1
; SEQ ID NO 6
; LENGTH: 24
; TYPE: PRT
; ORGANISM: Artificial sequence
; FEATURE:
; OTHER INFORMATION: Synthetic peptide
US-10-413-785-6

Query Match      29.9%; Score 29; DB 14; Length 24;
Best Local Similarity 41.7%; Pred. No. 9.9e+02;
Matches 5; Conservative 2; Mismatches 5; Indels 0; Gaps 0;

Qy      8 KIAFKIVSQEPA 19
Db      8 KVGFPVTFQVPA 19

RESULT 98
US-10-437-963-177203
; Sequence 177203, Application US/10437963
; Publication No. US20040123343A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa, Thomas J.
; APPLICANT: Kovalic, David K.
; APPLICANT: Zhou, Yihua
; APPLICANT: Cao, Yongwei
; APPLICANT: Wu, Wei
; APPLICANT: Boukharov, Andrey A.
; APPLICANT: Barbazuk, Brad
; APPLICANT: Li, Ping
; TITLE OF INVENTION: Rice Nucleic Acid Molecules and Other Molecules Associated With
; TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement
; FILE REFERENCE: 38-21(53221)B
; CURRENT APPLICATION NUMBER: US/10/437,963
; CURRENT FILING DATE: 2003-05-14
; NUMBER OF SEQ ID NOS: 204966
; SEQ ID NO 177203
; LENGTH: 26
; TYPE: PRT
; ORGANISM: Oryza sativa
; FEATURE:
; OTHER INFORMATION: Clone ID: PAT_MRT4530_74879C.1.pep
US-10-437-963-177203

Query Match      29.9%; Score 29; DB 16; Length 26;
Best Local Similarity 64.3%; Pred. No. 1.1e+03;
Matches 9; Conservative 0; Mismatches 3; Indels 2; Gaps 1;

Qy      4 HLNSKIAFKIVSQE 17
Db      6 HLES--AFKIFSIE 17

RESULT 99
US-09-864-761-40576
; Sequence 40576, Application US/09864761
; Patent No. US20020048763A1
; GENERAL INFORMATION:
; APPLICANT: Penn, Sharron G.
; APPLICANT: Rank, David R.
; APPLICANT: Hanzel, David K.
; APPLICANT: Chen, Wensheng
; TITLE OF INVENTION: HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES USEFUL FOR
; TITLE OF INVENTION: GENE EXPRESSION ANALYSIS BY MICROARRAY
; FILE REFERENCE: Aeomica-X-1
; CURRENT APPLICATION NUMBER: US/09/864,761
; CURRENT FILING DATE: 2001-05-23
```

; PRIOR APPLICATION NUMBER: US 60/180,312  
; PRIOR FILING DATE: 2000-02-04  
; PRIOR APPLICATION NUMBER: US 60/207,456  
; PRIOR FILING DATE: 2000-05-26  
; PRIOR APPLICATION NUMBER: US 09/632,366  
; PRIOR FILING DATE: 2000-08-03  
; PRIOR APPLICATION NUMBER: GB 24263.6  
; PRIOR FILING DATE: 2000-10-04  
; PRIOR APPLICATION NUMBER: US 60/236,359  
; PRIOR FILING DATE: 2000-09-27  
; PRIOR APPLICATION NUMBER: PCT/US01/00666  
; PRIOR FILING DATE: 2001-01-30  
; PRIOR APPLICATION NUMBER: PCT/US01/00667  
; PRIOR FILING DATE: 2001-01-30  
; PRIOR APPLICATION NUMBER: PCT/US01/00664  
; PRIOR FILING DATE: 2001-01-30  
; PRIOR APPLICATION NUMBER: PCT/US01/00669  
; PRIOR FILING DATE: 2001-01-30  
; PRIOR APPLICATION NUMBER: PCT/US01/00665  
; PRIOR FILING DATE: 2001-01-30  
; PRIOR APPLICATION NUMBER: PCT/US01/00668  
; PRIOR FILING DATE: 2001-01-30  
; PRIOR APPLICATION NUMBER: PCT/US01/00663  
; PRIOR FILING DATE: 2001-01-30  
; PRIOR APPLICATION NUMBER: PCT/US01/00662  
; PRIOR FILING DATE: 2001-01-30  
; PRIOR APPLICATION NUMBER: PCT/US01/00661  
; PRIOR FILING DATE: 2001-01-30  
; PRIOR APPLICATION NUMBER: PCT/US01/00670  
; PRIOR FILING DATE: 2001-01-30  
; PRIOR APPLICATION NUMBER: US 60/234,687  
; PRIOR FILING DATE: 2000-09-21  
; PRIOR APPLICATION NUMBER: US 09/608,408  
; PRIOR FILING DATE: 2000-06-30  
; PRIOR APPLICATION NUMBER: US 09/774,203  
; PRIOR FILING DATE: 2001-01-29  
; NUMBER OF SEQ ID NOS: 49117  
; SOFTWARE: Annomax Sequence Listing Engine vers. 1.1  
; SEQ ID NO 40576  
; LENGTH: 28  
; TYPE: PRT  
; ORGANISM: Homo sapiens  
; FEATURE:  
; OTHER INFORMATION: MAP TO AL109824.21  
; OTHER INFORMATION: EXPRESSED IN PLACENTA, SIGNAL = 1  
; OTHER INFORMATION: EXPRESSED IN FETAL LIVER, SIGNAL = 1.1  
; OTHER INFORMATION: EXPRESSED IN BONE MARROW, SIGNAL = 1.1  
; OTHER INFORMATION: EXPRESSED IN LUNG, SIGNAL = 0.83  
; OTHER INFORMATION: EXPRESSED IN ADULT LIVER, SIGNAL = 0.96  
; OTHER INFORMATION: EXPRESSED IN HEART, SIGNAL = 1.4  
; OTHER INFORMATION: EXPRESSED IN BRAIN, SIGNAL = 1  
US-09-864-761-40576

Query Match 29.9%; Score 29; DB 9; Length 28;  
Best Local Similarity 42.9%; Pred. No. 1.2e+03;  
Matches 6; Conservative 3; Mismatches 5; Indels 0; Gaps 0;

QY 5 LNSKIAFKIVSQEP 18  
|||:|:|:  
DB 5 LKSKASFSIYAFDP 18

RESULT 100  
US-09-982-172-128  
; Sequence 128, Application US/09982172  
; Patent No. US20020137119A1  
; GENERAL INFORMATION:  
; APPLICANT: Emil Israel Katz  
; TITLE OF INVENTION: PEPTIDES REPRESENTATIVE OF POLYPEPTIDES OF INTEREST AND ANTIBODIES  
; TITLE OF INVENTION: DIRECTED THEREAGAINST, AND METHODS, SYSTEMS AND KITS FOR GENERAT  
; TITLE OF INVENTION: UTILIZING EACH  
; FILE REFERENCE: 01/22283  
; CURRENT APPLICATION NUMBER: US/09/982,172

; CURRENT FILING DATE: 2001-10-19  
; NUMBER OF SEQ ID NOS: 253  
; SOFTWARE: PatentIn version 3.1  
; SEQ ID NO 128  
; LENGTH: 28  
; TYPE: PRT  
; ORGANISM: Artificial sequence  
; FEATURE:  
; OTHER INFORMATION: Computer generated synthetic peptide  
US-09-982-172-128

Query Match 29.9%; Score 29; DB 9; Length 28;  
Best Local Similarity 100.0%; Pred. No. 1.2e+03;  
Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 13 IVSQEP 18  
|||||  
DB 5 IVSQEP 10

## RESULT 101

US-10-029-386-29166  
; Sequence 29166, Application US/10029386  
; Publication No. US20030194704A1  
; GENERAL INFORMATION:  
; APPLICANT: Penn, Sharon G.  
; APPLICANT: Rank, David R.  
; APPLICANT: Hanzel, David K.  
; TITLE OF INVENTION: HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES USEFUL FOR  
; TITLE OF INVENTION: EXPRESSION ANALYSIS TWO  
; FILE REFERENCE: ACOMICA-X-2  
; CURRENT APPLICATION NUMBER: US/10/029,386  
; CURRENT FILING DATE: 2001-12-20  
; NUMBER OF SEQ ID NOS: 34288  
; SOFTWARE: Annomax Sequence Listing Engine vers. 1.1  
; SEQ ID NO 29166  
; LENGTH: 32  
; TYPE: PRT  
; ORGANISM: Homo sapiens  
; FEATURE:  
; OTHER INFORMATION: MAP TO CHR7.1  
; OTHER INFORMATION: EXPRESSED IN BONE MARROW, SIGNAL = 1.4  
; OTHER INFORMATION: EXPRESSED IN HEART, SIGNAL = 1.3  
; OTHER INFORMATION: EXPRESSED IN FETAL LIVER, SIGNAL = 1.2  
; OTHER INFORMATION: EXPRESSED IN LUNG, SIGNAL = 1.1  
; OTHER INFORMATION: EXPRESSED IN HELA, SIGNAL = 1.1  
; OTHER INFORMATION: EXPRESSED IN BRAIN, SIGNAL = 1.4  
US-10-029-386-29166

Query Match 29.9%; Score 29; DB 14; Length 32;  
Best Local Similarity 71.4%; Pred. No. 1.4e+03;  
Matches 5; Conservative 2; Mismatches 0; Indels 0; Gaps 0;

QY 12 KIVSQEP 18  
|:|:|:  
DB 17 KIVSQEP 23

## RESULT 102

US-09-764-891-4901  
; Sequence 4901, Application US/09764891  
; Publication No. US20030077808A1  
; GENERAL INFORMATION:  
; APPLICANT: Rosen et al.  
; TITLE OF INVENTION: Nucleic Acids, Proteins, and Antibodies  
; FILE REFERENCE: PC006  
; CURRENT APPLICATION NUMBER: US/09/764,891  
; CURRENT FILING DATE: 2001-01-17  
; Prior application data removed - consult PALM or file wrapper  
; NUMBER OF SEQ ID NOS: 10231  
; SOFTWARE: PatentIn Ver. 2.0  
; SEQ ID NO 4901  
; LENGTH: 33

```

; TYPE: PRT
; ORGANISM: Homo sapiens
; FEATURE:
; NAME/KEY: SITE
; LOCATION: (29)
; OTHER INFORMATION: Xaa equals any of the naturally occurring L-amino acids
US-09-764-891-4901

```

Query Match 29.9%; Score 29; DB 10; Length 33;

Best Local Similarity 50.0%; Pred. No. 1.4e+03;

Matches 5; Conservative 2; Mismatches 3; Indels 0; Gaps 0;

```

QY 2 PNHLNSKIAF 11
DB 14 PNKLTSQLTF 23

```

#### RESULT 103

```

US-10-091-414-161
; Sequence 161, Application US/10091414
; Publication No. US20030224461A1
; GENERAL INFORMATION:
; APPLICANT: Rosen et al.
; TITLE OF INVENTION: Nucleic Acids, Proteins, and Antibodies
; FILE REFERENCE: PAIL6C1
; CURRENT APPLICATION NUMBER: US/10/091,414
; CURRENT FILING DATE: 2002-03-07
; Prior Application removed - See File Wrapper or Palm
; NUMBER OF SEQ ID NOS: 392
; SOFTWARE: PatentIn ver. 2.0
; SEQ ID NO 161
; LENGTH: 33
; TYPE: PRT
; ORGANISM: Homo sapiens
; FEATURE:
; NAME/KEY: misc_feature
; LOCATION: (29)
; OTHER INFORMATION: Xaa equals any of the naturally occurring L-amino acids
US-10-091-414-161

```

Query Match 29.9%; Score 29; DB 14; Length 33;

Best Local Similarity 50.0%; Pred. No. 1.4e+03;

Matches 5; Conservative 2; Mismatches 3; Indels 0; Gaps 0;

```

QY 2 PNHLNSKIAF 11
DB 14 PNKLTSQLTF 23

```

#### RESULT 104

```

US-10-413-785-4
; Sequence 4, Application US/10413785
; Publication No. US20030229906A1
; GENERAL INFORMATION:
; APPLICANT: Gelman et al.
; TITLE OF INVENTION: METHODS AND COMPOSITIONS FOR THE TREATMENT OF DISORDERS OF HIV
; FILE REFERENCE: 29636/38269A
; CURRENT APPLICATION NUMBER: US/10/413,785
; CURRENT FILING DATE: 2003-04-14
; Prior Application Number: US 60/372,557
; Prior Filing Date: 2002-04-15
; NUMBER OF SEQ ID NOS: 15
; SOFTWARE: PatentIn version 3.1
; SEQ ID NO 4
; LENGTH: 33
; TYPE: PRT
; ORGANISM: Artificial sequence
; FEATURE:
; OTHER INFORMATION: Synthetic peptide
US-10-413-785-4

```

Query Match 29.9%; Score 29; DB 14; Length 33;

Best Local Similarity 41.7%; Pred. No. 1.4e+03;

Matches 5; Conservative 2; Mismatches 5; Indels 0; Gaps 0;

```

QY 8 KIAFKIVSQEPA 19
DB 17 KVGFEVTPQVPA 28

```

#### RESULT 105

```

US-10-351-641-1670
; Sequence 1670, Application US/10351641
; Publication No. US20030186874A1
; GENERAL INFORMATION:
; APPLICANT: Barney, S.
; APPLICANT: Guthrie, K.
; APPLICANT: Merutka, G.
; APPLICANT: Anwer, M.
; APPLICANT: Lambert, D.
; TITLE OF INVENTION: HYBRID POLYPEPTIDES WITH ENHANCED PHARMACOKINETIC
; TITLE OF INVENTION: PROPERTIES
; FILE REFERENCE: 7872-100
; CURRENT APPLICATION NUMBER: US/10/351,641
; CURRENT FILING DATE: 2003-01-24
; Prior Application Number: 09/350,641
; Prior Filing Date: 1999-07-09
; Prior Application Number: 09/315,304
; Prior Filing Date: 1999-05-20
; Prior Application Number: 09/082,279
; Prior Filing Date: 1998-05-20
; NUMBER OF SEQ ID NOS: 1757
; SOFTWARE: FastSeq for Windows Version 3.0
; SEQ ID NO 1670
; LENGTH: 34
; TYPE: PRT
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Core polypeptide
; NAME/KEY: SITE
; LOCATION: 17
; OTHER INFORMATION: Xaa = U (Aminobutyric Acid)
US-10-351-641-1670

```

Query Match 29.9%; Score 29; DB 14; Length 34;

Best Local Similarity 40.0%; Pred. No. 1.5e+03;

Matches 6; Conservative 2; Mismatches 7; Indels 0; Gaps 0;

```

QY 3 NHLNSKIAFKIVSQE 17
DB 15 NKXNGTDAVKLIKQE 29

```

#### RESULT 106

```

US-10-351-641-1671
; Sequence 1671, Application US/10351641
; Publication No. US20030186874A1
; GENERAL INFORMATION:
; APPLICANT: Barney, S.
; APPLICANT: Guthrie, K.
; APPLICANT: Merutka, G.
; APPLICANT: Anwer, M.
; APPLICANT: Lambert, D.
; TITLE OF INVENTION: HYBRID POLYPEPTIDES WITH ENHANCED PHARMACOKINETIC
; TITLE OF INVENTION: PROPERTIES
; FILE REFERENCE: 7872-100
; CURRENT APPLICATION NUMBER: US/10/351,641
; CURRENT FILING DATE: 2003-01-24
; Prior Application Number: 09/350,641
; Prior Filing Date: 1999-07-09
; Prior Application Number: 09/315,304
; Prior Filing Date: 1999-05-20
; Prior Application Number: 09/082,279
; Prior Filing Date: 1998-05-20

```

```
; NUMBER OF SEQ ID NOS: 1757
; SOFTWARE: FastSeq for Windows Version 3.0
; SEQ ID NO 1671
; LENGTH: 34
; TYPE: PRT
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Core polypeptide
; NAME/KEY: SITE
; LOCATION: 16
; OTHER INFORMATION: Xaa = U (Aminobutyric Acid)
US-10-351-641-1671

Query Match      29.9%; Score 29; DB 14; Length 34;
Best Local Similarity 40.0%; Pred. No. 1.5e+03;
Matches 6; Conservative 2; Mismatches 7; Indels 0; Gaps 0;

QY 3 NHLNSKIAPKIVSQE 17
Db 14 NKXNGTDAVKLIKQE 28

RESULT 107
US-10-351-641-1672
; Sequence 1672, Application US/10351641
; Publication No. US20030186874A1
; GENERAL INFORMATION:
; APPLICANT: Barney, S.
; APPLICANT: Guthrie, K.
; APPLICANT: Merutka, G.
; APPLICANT: Anwer, M.
; APPLICANT: Lambert, D.
; TITLE OF INVENTION: HYBRID POLYPEPTIDES WITH ENHANCED PHARMACOKINETIC
; FILE REFERENCE: 7872-100
; CURRENT APPLICATION NUMBER: US/10/351,641
; PRIOR FILING DATE: 2003-01-24
; PRIOR APPLICATION NUMBER: 09/350,641
; PRIOR FILING DATE: 1998-07-09
; PRIOR APPLICATION NUMBER: 09/315,304
; PRIOR FILING DATE: 1998-05-20
; PRIOR APPLICATION NUMBER: 09/082,279
; PRIOR FILING DATE: 1998-05-20
; NUMBER OF SEQ ID NOS: 1757
; SOFTWARE: FastSeq for Windows Version 3.0
; SEQ ID NO 1672
; LENGTH: 34
; TYPE: PRT
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Core polypeptide
; NAME/KEY: SITE
; LOCATION: 15
; OTHER INFORMATION: Xaa = U (Aminobutyric Acid)
US-10-351-641-1672

Query Match      29.9%; Score 29; DB 14; Length 34;
Best Local Similarity 40.0%; Pred. No. 1.5e+03;
Matches 6; Conservative 2; Mismatches 7; Indels 0; Gaps 0;

QY 3 NHLNSKIAPKIVSQE 17
Db 13 NKXNGTDAVKLIKQE 27

RESULT 108
US-10-437-963-179203
; Sequence 179203, Application US/10437963
; Publication No. US20040123343A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa, Thomas J.
```

```
; APPLICANT: Kovalic, David K.
; APPLICANT: Zhou, Yihua
; APPLICANT: Cao, Yongwei
; APPLICANT: Wu, Wei
; APPLICANT: Boukharov, Andrey A.
; APPLICANT: Barbazuk, Brad
; APPLICANT: Li, Ping
; TITLE OF INVENTION: Rice Nucleic Acid Molecules and Other Molecules Associated With
; FILE REFERENCE: 38-21(53221)B
; CURRENT APPLICATION NUMBER: US/10/437,963
; CURRENT FILING DATE: 2003-05-14
; NUMBER OF SEQ ID NOS: 204966
; SEQ ID NO 179203
; LENGTH: 34
; TYPE: PRT
; ORGANISM: Oryza sativa
; FEATURE:
; NAME/KEY: unsure
; LOCATION: (1)..(34)
; OTHER INFORMATION: unsure at all Xaa locations
; FEATURE:
; OTHER INFORMATION: Clone ID: PAT_MRT4530_7668C.1.pap
US-10-437-963-179203

Query Match      29.9%; Score 29; DB 16; Length 34;
Best Local Similarity 45.5%; Pred. No. 1.5e+03;
Matches 5; Conservative 2; Mismatches 4; Indels 0; Gaps 0;

QY 4 HLNSKIAPKIV 14
Db 24 HITSXITPPII 34

RESULT 109
US-09-820-649-195
; Sequence 195, Application US/09820649
; Publication No. US20030199683A1
; GENERAL INFORMATION:
; APPLICANT: Ruben et al.
; TITLE OF INVENTION: 83 Human Secreted Proteins
; FILE REFERENCE: PZO12PI
; CURRENT APPLICATION NUMBER: US/09/820,649
; CURRENT FILING DATE: 2001-03-30
; PRIOR APPLICATION NUMBER: US/09/236,557
; PRIOR FILING DATE: 1999-01-26
; PRIOR APPLICATION NUMBER: PCT/US98/15949
; PRIOR FILING DATE: 1998-07-29
; PRIOR APPLICATION NUMBER: 60/054,212
; PRIOR FILING DATE: 1997-07-30
; PRIOR APPLICATION NUMBER: 60/054,209
; PRIOR FILING DATE: 1997-07-30
; PRIOR APPLICATION NUMBER: 60/054,234
; PRIOR FILING DATE: 1997-07-30
; PRIOR APPLICATION NUMBER: 60/054,218
; PRIOR FILING DATE: 1997-07-30
; PRIOR APPLICATION NUMBER: 60/054,214
; PRIOR FILING DATE: 1997-07-30
; PRIOR APPLICATION NUMBER: 60/054,236
; PRIOR FILING DATE: 1997-07-30
; PRIOR APPLICATION NUMBER: 60/054,215
; PRIOR FILING DATE: 1997-07-30
; PRIOR APPLICATION NUMBER: 60/054,211
; Remaining Prior Application data removed - See File Wrapper or PALM.
; NUMBER OF SEQ ID NOS: 353
; SOFTWARE: Patentin Ver. 2.0
; SEQ ID NO 195
; LENGTH: 37
; TYPE: PRT
; ORGANISM: Homo sapiens
US-09-820-649-195
```



Query Match 29.9%; Score 29; DB 10; Length 37;  
Best Local Similarity 54.5%; Pred. No. 1.6e+03;  
Matches 6; Conservative 1; Mismatches 4; Indels 0; Gaps 0;

QY 3 NHLNSKIAFKI 13  
: || | | |  
Db 11 HHLKSKPHLKI 21

## RESULT 110

US-10-160-162-195  
; Sequence 195, Application US/10160162  
; Publication No. US20030166541A1  
; GENERAL INFORMATION:  
; APPLICANT: Ruben et al.  
; TITLE OF INVENTION: 83 Human Secreted Proteins  
; FILE REFERENCE: P2012P2  
; CURRENT APPLICATION NUMBER: US/10/160,162  
; CURRENT FILING DATE: 2002-06-04  
; PRIOR APPLICATION NUMBER: 60/295,558  
; PRIOR FILING DATE: 2001-06-05  
; PRIOR APPLICATION NUMBER: 09/236,557  
; PRIOR FILING DATE: 1999-01-26  
; PRIOR APPLICATION NUMBER: PCT/US98/15949  
; PRIOR FILING DATE: 1998-07-29  
; PRIOR APPLICATION NUMBER: 60/054,212  
; PRIOR FILING DATE: 1997-07-30  
; PRIOR APPLICATION NUMBER: 60/054,209  
; PRIOR FILING DATE: 1997-07-30  
; PRIOR APPLICATION NUMBER: 60/054,214  
; PRIOR FILING DATE: 1997-07-30  
; PRIOR APPLICATION NUMBER: 60/054,236  
; PRIOR FILING DATE: 1997-07-30  
; PRIOR APPLICATION NUMBER: 60/054,215  
; PRIOR FILING DATE: 1997-07-30  
; PRIOR APPLICATION NUMBER: 60/054,211  
; PRIOR FILING DATE: 1997-07-30  
; PRIOR APPLICATION NUMBER: 60/054,217  
; PRIOR FILING DATE: 1997-07-30  
; PRIOR APPLICATION NUMBER: 60/054,213  
; PRIOR FILING DATE: 1997-07-30  
; PRIOR APPLICATION NUMBER: 60/055,968  
; PRIOR FILING DATE: 1997-08-18  
; PRIOR APPLICATION NUMBER: 60/055,969  
; PRIOR FILING DATE: 1997-08-18  
; PRIOR APPLICATION NUMBER: 60/055,972  
; PRIOR FILING DATE: 1997-08-18  
; PRIOR APPLICATION NUMBER: 60/056,561  
; PRIOR FILING DATE: 1997-08-19  
; PRIOR APPLICATION NUMBER: 60/056,534  
; PRIOR FILING DATE: 1997-08-19  
; PRIOR APPLICATION NUMBER: 60/056,729  
; PRIOR FILING DATE: 1997-08-19  
; PRIOR APPLICATION NUMBER: 60/056,543  
; PRIOR FILING DATE: 1997-08-19  
; PRIOR APPLICATION NUMBER: 60/056,727  
; PRIOR FILING DATE: 1997-08-19  
; PRIOR APPLICATION NUMBER: 60/056,554  
; PRIOR FILING DATE: 1997-08-19  
; PRIOR APPLICATION NUMBER: 60/056,730  
; PRIOR FILING DATE: 1997-08-19  
; NUMBER OF SEQ ID NOS: 353  
; SOFTWARE: Patent In Ver. 2.0  
; SEQ ID NO 195  
; LENGTH: 37  
; TYPE: PRT  
; ORGANISM: Homo sapiens  
US-10-160-162-195

Query Match 29.9%; Score 29; DB 14; Length 37;  
Best Local Similarity 54.5%; Pred. No. 1.6e+03;  
Matches 6; Conservative 1; Mismatches 4; Indels 0; Gaps 0;

QY 3 NHLNSKIAFKI 13  
: || | | |  
Db 11 HHLKSKPHLKI 21

## RESULT 111

US-10-437-963-107393  
; Sequence 107393, Application US/10437963  
; Publication No. US20040123343A1  
; GENERAL INFORMATION:  
; APPLICANT: La Rosa, Thomas J.  
; APPLICANT: Kovalic, David K.  
; APPLICANT: Zhou, Yihua  
; APPLICANT: Cao, Yongwei  
; APPLICANT: Wu, Wei  
; APPLICANT: Boukharov, Andrey A.  
; APPLICANT: Barbazuk, Brad  
; APPLICANT: Li, Ping  
; TITLE OF INVENTION: Rice Nucleic Acid Molecules and Other Molecules Associated With  
; FILE REFERENCE: 38-21(53221)B  
; CURRENT APPLICATION NUMBER: US/10/437,963  
; CURRENT FILING DATE: 2003-05-14  
; NUMBER OF SEQ ID NOS: 204966  
; SEQ ID NO 107393  
; LENGTH: 37  
; TYPE: PRT  
; ORGANISM: Oryza sativa  
; FEATURE:  
; OTHER INFORMATION: Clone ID: PAT\_MRT4530\_1174C.1.pep  
US-10-437-963-107393

Query Match 29.9%; Score 29; DB 16; Length 37;  
Best Local Similarity 80.0%; Pred. No. 1.6e+03;  
Matches 4; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 2 PNHLN 6  
: || | |  
Db 25 PNHLN 29

## RESULT 112

US-09-250-883-21  
; Sequence 21, Application US/09250883  
; Patent No. US20020042049A1  
; GENERAL INFORMATION:  
; APPLICANT: Russell, John  
; APPLICANT: Colpitts, Tracey  
; TITLE OF INVENTION: REAGENTS AND METHODS USEFUL  
; TITLE OF INVENTION: FOR DETECTING DISEASE OF THE BREAST  
; NUMBER OF SEQUENCES: 23  
; CORRESPONDENCE ADDRESS:  
; ADDRESSEE: Abbott Laboratories  
; STREET: 100 Abbott Park Road  
; CITY: Abbott Park  
; STATE: IL  
; COUNTRY: USA  
; ZIP: 60064-3500  
; COMPUTER READABLE FORM:  
; MEDIUM TYPE: Diskette  
; COMPUTER: IBM Compatible  
; OPERATING SYSTEM: DOS  
; SOFTWARE: FastSeq for Windows Version 2.0  
; CURRENT APPLICATION DATA:  
; APPLICATION NUMBER: US/09/250,883  
; FILING DATE:  
; CLASSIFICATION:  
; PRIOR APPLICATION DATA:  
; APPLICATION NUMBER: US/08/889,316

```

; FILING DATE:
; ATTORNEY/AGENT INFORMATION:
; NAME: Becker, Cheryl L
; REGISTRATION NUMBER: 35,441
; REFERENCE/DOCKET NUMBER: 6131.US.01
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: 847/935-1729
; TELEFAX: 847/938-2623
; TELEX:
; INFORMATION FOR SEQ ID NO: 21:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 38 amino acids
; TYPE: amino acid
; STRANDEDNESS: single
; TOPOLOGY: linear
; MOLECULE TYPE: No. US20020042049A1e
US-09-250-883-21

Query Match 29.9%; Score 29; DB 9; Length 38;
Best Local Similarity 83.3%; Pred. No. 1.7e+03;
Matches 5; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 4 HLNSKI 9
Db 18 HLNSKL 23

RESULT 113
US-09-925-299-1490
; Sequence 1490, Application US/09925299
; Patent No. US20020055627A1
; GENERAL INFORMATION:
; APPLICANT: Rosen et al.
; TITLE OF INVENTION: Nucleic Acids, Proteins and Antibodies
; FILE REFERENCE: PA102
; CURRENT APPLICATION NUMBER: US/09/925,299
; CURRENT FILING DATE: 2001-08-10
; PRIOR APPLICATION NUMBER: PCT/US00/05883
; PRIOR FILING DATE: 2000-03-08
; PRIOR APPLICATION NUMBER: 60/124,270
; PRIOR FILING DATE: 1999-03-12
; NUMBER OF SEQ ID NOS: 1556
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 1490
; LENGTH: 39
; TYPE: PRT
; ORGANISM: Homo sapiens
; FEATURE:
; NAME/KEY: SITE
; LOCATION: (5)
; OTHER INFORMATION: Xaa equals any of the naturally occurring L-amino acids
; NAME/KEY: SITE
; LOCATION: (8)
; OTHER INFORMATION: Xaa equals any of the naturally occurring L-amino acids
; NAME/KEY: SITE
; LOCATION: (12)
; OTHER INFORMATION: Xaa equals any of the naturally occurring L-amino acids
; NAME/KEY: SITE
; LOCATION: (28)
; OTHER INFORMATION: Xaa equals any of the naturally occurring L-amino acids
; NAME/KEY: SITE
; LOCATION: (35)
; OTHER INFORMATION: Xaa equals any of the naturally occurring L-amino acids
; NAME/KEY: SITE
; LOCATION: (37)
; OTHER INFORMATION: Xaa equals any of the naturally occurring L-amino acids
US-09-925-299-1490

Query Match 29.9%; Score 29; DB 9; Length 39;
Best Local Similarity 37.5%; Pred. No. 1.7e+03;
Matches 6; Conservative 3; Mismatches 7; Indels 0; Gaps 0;

QY 3 NHLNSKIAPKIVSQEP 18
```

```

Db 14 NLMTGRHSFKTYSQXP 29

RESULT 114
US-09-071-838-175
; Sequence 175, Application US/09071838
; Patent No. US20020152501A1
; GENERAL INFORMATION:
; APPLICANT: Fischer, Robert L.
; APPLICANT: Chad, Nir
; APPLICANT: Kiyosue, Tomohiro
; APPLICANT: Yadegari, Ramin
; APPLICANT: Margossian, Linda
; APPLICANT: Harada, John
; APPLICANT: Goldberg, Robert B.
; TITLE OF INVENTION: Nucleic Acids That Control Seed and
; TITLE OF INVENTION: Fruit Development in Plants
; NUMBER OF SEQUENCES: 324
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Townsend and Townsend and Crew LLP
; STREET: Two Embarcadero Center, Eighth Floor
; CITY: San Francisco
; STATE: California
; COUNTRY: USA
; ZIP: 94111-3834
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: PatentIn Release #1.0, Version #1.30
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/09/071,838
; FILING DATE: 01-MAY-1998
; CLASSIFICATION: 800
; ATTORNEY/AGENT INFORMATION:
; NAME: Bastian, Kevin L.
; REGISTRATION NUMBER: 34,774
; REFERENCE/DOCKET NUMBER: 023070-086100US
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (415) 576-0200
; TELEFAX: (415) 576-0300
; INFORMATION FOR SEQ ID NO: 175:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 39 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
US-09-071-838-175

Query Match 29.9%; Score 29; DB 9; Length 39;
Best Local Similarity 33.3%; Pred. No. 1.7e+03;
Matches 4; Conservative 5; Mismatches 3; Indels 0; Gaps 0;

QY 3 NHLNSKIAPKIV 14
Db 19 NHVNIRISLIVI 30

RESULT 115
US-09-925-299-1490
; Sequence 1490, Application US/09925299
; Publication No. US20030040617A9
; GENERAL INFORMATION:
; APPLICANT: Rosen et al.
; TITLE OF INVENTION: Nucleic Acids, Proteins and Antibodies
; FILE REFERENCE: PA102
; CURRENT APPLICATION NUMBER: US/09/925,299
; CURRENT FILING DATE: 2001-08-10
; PRIOR APPLICATION NUMBER: PCT/US00/05883
; PRIOR FILING DATE: 2000-03-08
; PRIOR APPLICATION NUMBER: 60/124,270
; PRIOR FILING DATE: 1999-03-12
```



## RESULT 119

US-10-029-386-33231

; Sequence 33231, Application US/10029386

; Publication No. US20030194704A1

; GENERAL INFORMATION:

; APPLICANT: Penn, Sharon G.

; APPLICANT: Rank, David R.

; APPLICANT: Hanzel, David K.

; TITLE OF INVENTION: HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES USEFUL FOR G

; FILE REFERENCE: AEWICA-X-2

; CURRENT APPLICATION NUMBER: US/10/029,386

; CURRENT FILING DATE: 2001-12-20

; NUMBER OF SEQ ID NOS: 34288

; SOFTWARE: Anomax Sequence Listing Engine vers. 1.1

; SEQ ID NO 33231

; LENGTH: 41

; TYPE: PRT

; ORGANISM: Homo sapiens

; FEATURE:

; OTHER INFORMATION: MAP TO AC022127.3

; OTHER INFORMATION: EXPRESSED IN BRAIN, SIGNAL = 3.6

; OTHER INFORMATION: EXPRESSED IN PLACENTA, SIGNAL = 3.5

; OTHER INFORMATION: EXPRESSED IN LUNG, SIGNAL = 4.7

; OTHER INFORMATION: EXPRESSED IN HELA, SIGNAL = 4.1

; OTHER INFORMATION: EXPRESSED IN FETAL LIVER, SIGNAL = 3.5

US-10-029-386-33231

Query Match

Best Local Similarity 29.9%; Score 29; DB 14; Length 41;

Matches 5; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 4 HLNSKI 9

Db 35 HLNSKL 40

|||||

|||||

## RESULT 120

US-10-437-963-140191

; Sequence 140191, Application US/10437963

; Publication No. US20040123343A1

; GENERAL INFORMATION:

; APPLICANT: La Rosa, Thomas J.

; APPLICANT: Kovalic, David K.

; APPLICANT: Zhou, Yihua

; APPLICANT: Cao, Yongwei

; APPLICANT: Wu, Wei

; APPLICANT: Boukharov, Andrey A.

; APPLICANT: Barbazuk, Brad

; APPLICANT: Li, Ping

; TITLE OF INVENTION: Rice Nucleic Acid Molecules and Other Molecules Associated With

; FILE REFERENCE: 38-21(53221)B

; CURRENT APPLICATION NUMBER: US/10/437,963

; CURRENT FILING DATE: 2003-05-14

; NUMBER OF SEQ ID NOS: 204966

; SEQ ID NO 140191

; LENGTH: 41

; TYPE: PRT

; ORGANISM: Oryza sativa

; FEATURE:

; OTHER INFORMATION: Clone ID: PAT\_MRT4530\_41412C.1.pap

US-10-437-963-140191

Query Match

Best Local Similarity 29.9%; Score 29; DB 16; Length 41;

Matches 5; Conservative 5; Mismatches 6; Indels 0; Gaps 0;

QY 1 EPNHLSKIAFKIVSQ 16

Db 13 EKKKINKKEIYLVNE 28

|||||

|||||

## RESULT 121

US-10-424-599-238139

; Sequence 238139, Application US/10424599

; Publication No. US20040031072A1

; GENERAL INFORMATION:

; APPLICANT: La Rosa, Thomas J

; APPLICANT: Kovalic, David K

; APPLICANT: Zhou Yihua

; APPLICANT: Cao Yongwei

; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With

; FILE REFERENCE: 38-21(53223)B

; CURRENT APPLICATION NUMBER: US/10/424,599

; CURRENT FILING DATE: 2003-04-28

; NUMBER OF SEQ ID NOS: 285684

; SEQ ID NO 238139

; LENGTH: 42

; TYPE: PRT

; ORGANISM: Glycine max

; FEATURE:

; OTHER INFORMATION: Clone ID: PAT\_MRT3847\_57065C.1.pap

US-10-424-599-238139

Query Match

Best Local Similarity 29.9%; Score 29; DB 15; Length 42;

Matches 3; Conservative 7; Mismatches 5; Indels 0; Gaps 0;

QY 1 EPNHLSKIAFKIVS 15

Db 17 DPNLNLQRLTYQFAT 31

|||||

|||||

## RESULT 122

US-10-424-599-271865

; Sequence 271865, Application US/10424599

; Publication No. US20040031072A1

; GENERAL INFORMATION:

; APPLICANT: La Rosa, Thomas J

; APPLICANT: Kovalic, David K

; APPLICANT: Zhou Yihua

; APPLICANT: Cao Yongwei

; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With

; FILE REFERENCE: 38-21(53223)B

; CURRENT APPLICATION NUMBER: US/10/424,599

; CURRENT FILING DATE: 2003-04-28

; NUMBER OF SEQ ID NOS: 285684

; SEQ ID NO 271865

; LENGTH: 42

; TYPE: PRT

; ORGANISM: Glycine max

; FEATURE:

; OTHER INFORMATION: Clone ID: PAT\_MRT3847\_87512C.1.pap

US-10-424-599-271865

Query Match

Best Local Similarity 29.9%; Score 29; DB 15; Length 42;

Matches 7; Conservative 4; Mismatches 6; Indels 4; Gaps 1;

QY 2 FNH----LNSKIAFKIVSQEP 18

Db 1 PSHKTFRIKKKLAKKIKQNK 21

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|||||

## RESULT 123

US-10-437-963-151559

; Sequence 151559, Application US/10437963

; Publication No. US20040123343A1

; GENERAL INFORMATION:

; APPLICANT: La Rosa, Thomas J.

; APPLICANT: Kovalic, David K.

; APPLICANT: Zhou, Yihua

```
/ APPLICANT: Cao, Yongwei
/ APPLICANT: Wu, Wei
/ APPLICANT: Boukharov, Andrey A.
/ APPLICANT: Barbazuk, Brad
/ APPLICANT: Li, Ping
/ TITLE OF INVENTION: Rice Nucleic Acid Molecules and Other Molecules Associated With
/ FILE REFERENCE: 38-21(53221)B
/ CURRENT APPLICATION NUMBER: US/10/437,963
/ CURRENT FILING DATE: 2003-05-14
/ NUMBER OF SEQ ID NOS: 204966
/ SEQ ID NO 151559
/ LENGTH: 43
/ TYPE: PRT
/ ORGANISM: Oryza sativa
/ FEATURE:
/ OTHER INFORMATION: Clone ID: PAT_MRT4530_51691C.1.pep
US-10-437-963-151559

Query Match          29.9%; Score 29; DB 16; Length 43;
Best Local Similarity 62.5%; Pred. No. 1.9e+03;
Matches 5; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

Qy 2 PNLNSKI 9
   ||:||||
Db 29 PSYLGKI 36

RESULT 124
US-09-989-919-94
/ Sequence 94, Application US/09989919
/ Patent No. US20020164344A1
/ GENERAL INFORMATION:
/ APPLICANT: Macina, Roberto
/ APPLICANT: Recipon, Herve
/ APPLICANT: Pluta, Jason
/ APPLICANT: Ghosh, Malavika
/ APPLICANT: Sun, Yongming
/ APPLICANT: Liu, Chenghua
/ TITLE OF INVENTION: Compositions and Methods Relating to Colon Specific Genes and Pro
/ FILE REFERENCE: DEX-0289
/ CURRENT APPLICATION NUMBER: US/09/989,919
/ CURRENT FILING DATE: 2001-11-21
/ PRIOR APPLICATION NUMBER: 60/252,505
/ PRIOR FILING DATE: 2000-11-22
/ NUMBER OF SEQ ID NOS: 124
/ SOFTWARE: PatentIn version 3.1
/ SEQ ID NO 94
/ LENGTH: 44
/ TYPE: PRT
/ ORGANISM: Homo sapien
US-09-989-919-94

Query Match          29.9%; Score 29; DB 9; Length 44;
Best Local Similarity 38.5%; Pred. No. 2e+03;
Matches 5; Conservative 4; Mismatches 4; Indels 0; Gaps 0;

Qy 3 NHLNSKIAPKIVS 15
   ||:||:||
Db 18 NTLTKIKYLSIS 30

RESULT 125
US-10-424-599-221932
/ Sequence 221932, Application US/10424599
/ Publication No. US20040031072A1
/ GENERAL INFORMATION:
/ APPLICANT: La Rosa Thomas J
/ APPLICANT: Kovalic David K
/ APPLICANT: Zhou Yihua
/ APPLICANT: Cao Yongwei
/ TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With
```

```
/ FILE REFERENCE: 38-21(53223)B
/ CURRENT APPLICATION NUMBER: US/10/424,599
/ CURRENT FILING DATE: 2003-04-28
/ NUMBER OF SEQ ID NOS: 285684
/ SEQ ID NO 221932
/ LENGTH: 44
/ TYPE: PRT
/ ORGANISM: Glycine max
/ FEATURE:
/ OTHER INFORMATION: Clone ID: PAT_MRT3847_42434C.1.pep
US-10-424-599-221932

Query Match          29.9%; Score 29; DB 15; Length 44;
Best Local Similarity 62.5%; Pred. No. 2e+03;
Matches 5; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

Qy 1 EPNHLSNK 8
   ||:||
Db 37 EPPHLHSR 44

RESULT 126
US-10-437-963-124421
/ Sequence 124421, Application US/10437963
/ Publication No. US20040123343A1
/ GENERAL INFORMATION:
/ APPLICANT: La Rosa, Thomas J.
/ APPLICANT: Kovalic, David K.
/ APPLICANT: Zhou, Yihua
/ APPLICANT: Cao, Yongwei
/ APPLICANT: Wu, Wei
/ APPLICANT: Boukharov, Andrey A.
/ APPLICANT: Barbazuk, Brad
/ APPLICANT: Li, Ping
/ TITLE OF INVENTION: Rice Nucleic Acid Molecules and Other Molecules Associated With
/ FILE REFERENCE: 38-21(53221)B
/ CURRENT APPLICATION NUMBER: US/10/437,963
/ CURRENT FILING DATE: 2003-05-14
/ NUMBER OF SEQ ID NOS: 204966
/ SEQ ID NO 124421
/ LENGTH: 45
/ TYPE: PRT
/ ORGANISM: Oryza sativa
/ FEATURE:
/ OTHER INFORMATION: Clone ID: PAT_MRT4530_27161C.1.pep
US-10-437-963-124421

Query Match          29.9%; Score 29; DB 16; Length 45;
Best Local Similarity 62.5%; Pred. No. 2e+03;
Matches 5; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

Qy 3 NHLNSKIA 10
   ||:||
Db 37 NHFSTKIA 44

RESULT 127
US-09-933-767-340
/ Sequence 340, Application US/099333767
/ Publication No. US20030181692A1
/ GENERAL INFORMATION:
/ APPLICANT: Ni et al.
/ TITLE OF INVENTION: 207 Human Secreted Proteins
/ FILE REFERENCE: P2007P2
/ CURRENT APPLICATION NUMBER: US/09/933,767
/ CURRENT FILING DATE: 2001-08-22
/ PRIOR APPLICATION NUMBER: PCT/US01/05614
/ PRIOR FILING DATE: 2001-02-21
/ PRIOR APPLICATION NUMBER: 60/184,836
/ PRIOR FILING DATE: 2000-02-24
/ PRIOR APPLICATION NUMBER: 60/193,170
/ PRIOR FILING DATE: 2000-03-29
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Db 26 NLFTSQIKYIYSEKP 41

RESULT 129

US-10-023-282-340

; Sequence 340, Application US/10023282

; Publication No. US20030092893A1

; GENERAL INFORMATION:

; APPLICANT: Young et al.

; TITLE OF INVENTION: 207 Human Secreted Proteins

; FILE REFERENCE: PZ007P1

; CURRENT APPLICATION NUMBER: US/10/023,282

; CURRENT FILING DATE: 2001-12-20

; EARLIER APPLICATION NUMBER: 09/205,258

; EARLIER FILING DATE: 1998-12-04

; EARLIER APPLICATION NUMBER: PCT/US98/11422

; EARLIER FILING DATE: 1998-06-04

; EARLIER APPLICATION NUMBER: 60/048,885

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/049,375

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,881

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,880

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,896

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/049,020

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,876

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,895

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,884

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,894

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,971

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,964

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,882

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,899

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,893

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,900

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,901

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,892

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,915

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/049,019

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,970

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,972

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,916

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/049,373

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,875

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/049,374

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,917

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,949

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,974

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,883

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,897

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,898

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,962

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,963

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,877

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/048,878

; EARLIER FILING DATE: 1997-06-06

; EARLIER APPLICATION NUMBER: 60/070,923

; EARLIER FILING DATE: 1997-12-18

; EARLIER APPLICATION NUMBER: 60/092,921

; EARLIER FILING DATE: 1998-07-15

; EARLIER APPLICATION NUMBER: 60/094,657

; EARLIER FILING DATE: 1998-07-30

; NUMBER OF SEQ ID NOS: 1227

; SOFTWARE: PatentIn Ver. 2.0

; SEQ ID NO 340

; LENGTH: 46

; TYPE: PRT

; ORGANISM: Homo sapiens

US-10-023-282-340

Query Match 29.9%; Score 29; DB 14; Length 46;

Best Local Similarity 37.5%; Pred. No. 2.1e+03;

Matches 6; Conservative 4; Mismatches 6; Indels 0; Gaps 0;

Qy 3 NHLNSKIAFKIVSQEP 18

| : : : : |

| : : : : |

Db 26 NLFTSQIKYIYSEKP 41

RESULT 130

US-10-424-599-143969

; Sequence 143969, Application US/10424599

; Publication No. US20040031072A1

; GENERAL INFORMATION:

; APPLICANT: La Rosa, Thomas J

; APPLICANT: Kovalic, David K

; APPLICANT: Zhou, Yihua

; APPLICANT: Cao, Yongwei

; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With

; TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement

; FILE REFERENCE: 38-21(53223)B

; CURRENT APPLICATION NUMBER: US/10/424,599

; CURRENT FILING DATE: 2003-04-28

; NUMBER OF SEQ ID NOS: 285684

; SEQ ID NO 143969

; LENGTH: 46

; TYPE: PRT

; ORGANISM: Glycine max

; FEATURE:

; OTHER INFORMATION: Clone ID: PAT\_MRT3847\_101016C.1.pap

US-10-424-599-143969

Query Match 29.9%; Score 29; DB 15; Length 46;

Best Local Similarity 50.0%; Pred. No. 2.1e+03;

Matches 5; Conservative 2; Mismatches 3; Indels 0; Gaps 0;

Qy 4 HLNSKIAFKI 13

| : : : : |

| : : : : |

Db 33 HLNTSLNFSI 42

RESULT 131

US-10-424-599-194850

```
; Sequence 194850, Application US/10424599
; Publication No. US20040031072A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa Thomas J
; APPLICANT: Kovalic David K
; APPLICANT: Zhou Yihua
; APPLICANT: Cao Yongwei
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With
; TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement
; FILE REFERENCE: 38-21(53223)B
; CURRENT APPLICATION NUMBER: US/10/424,599
; CURRENT FILING DATE: 2003-04-28
; NUMBER OF SEQ ID NOS: 285684
; SEQ ID NO 194850
; LENGTH: 46
; TYPE: PRT
; ORGANISM: Glycine max
; FEATURE:
; OTHER INFORMATION: Clone ID: PAT_MRT3847_17978C.1.pap
US-10-424-599-194850

Query Match      29.9%; Score 29; DB 15; Length 46;
Best Local Similarity 38.5%; Pred. No. 2.1e+03;
Matches 5; Conservative 3; Mismatches 5; Indels 0; Gaps 0;

QY      2 PNHLNSKIAFKIV 14
DB      10 PYHLQTFHLKII 22

RESULT 132
US-10-424-599-196352
; Sequence 196352, Application US/10424599
; Publication No. US20040031072A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa Thomas J
; APPLICANT: Kovalic David K
; APPLICANT: Zhou Yihua
; APPLICANT: Cao Yongwei
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With
; TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement
; FILE REFERENCE: 38-21(53223)B
; CURRENT APPLICATION NUMBER: US/10/424,599
; CURRENT FILING DATE: 2003-04-28
; NUMBER OF SEQ ID NOS: 285684
; SEQ ID NO 196352
; LENGTH: 46
; TYPE: PRT
; ORGANISM: Glycine max
; FEATURE:
; OTHER INFORMATION: Clone ID: PAT_MRT3847_19332C.1.pap
US-10-424-599-196352

Query Match      29.9%; Score 29; DB 15; Length 46;
Best Local Similarity 71.4%; Pred. No. 2.1e+03;
Matches 5; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

QY      2 PNHLNSK 8
DB      35 PSHLTSK 41

RESULT 133
US-10-029-386-28419
; Sequence 28419, Application US/10029386
; Publication No. US20030194704A1
; GENERAL INFORMATION:
; APPLICANT: Penn, Sharon G.
; APPLICANT: Rank, David R.
; APPLICANT: Hanzel, David K.
; TITLE OF INVENTION: HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES USEFUL FOR C
; TITLE OF INVENTION: EXPRESSION ANALYSIS TWO
; FILE REFERENCE: AEOMICA-X-2
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; CURRENT APPLICATION NUMBER: US/10/029,386
; CURRENT FILING DATE: 2001-12-20
; NUMBER OF SEQ ID NOS: 34288
; SOFTWARE: Annomax Sequence Listing Engine vers. 1.1
; SEQ ID NO 28419
; LENGTH: 47
; TYPE: PRT
; ORGANISM: Homo sapiens
; FEATURE:
; OTHER INFORMATION: MAP TO CHR5.1
; OTHER INFORMATION: EXPRESSED IN LUNG, SIGNAL = 1.3
; OTHER INFORMATION: EXPRESSED IN HELA, SIGNAL = 1.6
; OTHER INFORMATION: EXPRESSED IN BRAIN, SIGNAL = 1.9
; OTHER INFORMATION: EXPRESSED IN HEART, SIGNAL = 1.8
; OTHER INFORMATION: EXPRESSED IN ADULT LIVER, SIGNAL = 2.8
; OTHER INFORMATION: EXPRESSED IN FETAL LIVER, SIGNAL = 1.4
; OTHER INFORMATION: SWISSPROT HIT: O27179, EVALUE 8.30e+00
US-10-029-386-28419

Query Match      29.9%; Score 29; DB 14; Length 47;
Best Local Similarity 44.4%; Pred. No. 2.1e+03;
Matches 4; Conservative 4; Mismatches 1; Indels 0; Gaps 0;

QY      2 PNHLNSKTA 10
DB      38 PSHLKSEVS 46

RESULT 134
US-10-424-599-162168
; Sequence 162168, Application US/10424599
; Publication No. US20040031072A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa Thomas J
; APPLICANT: Kovalic David K
; APPLICANT: Zhou Yihua
; APPLICANT: Cao Yongwei
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With
; TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement
; FILE REFERENCE: 38-21(53223)B
; CURRENT APPLICATION NUMBER: US/10/424,599
; CURRENT FILING DATE: 2003-04-28
; NUMBER OF SEQ ID NOS: 285684
; SEQ ID NO 162168
; LENGTH: 47
; TYPE: PRT
; ORGANISM: Glycine max
; FEATURE:
; OTHER INFORMATION: Clone ID: PAT_MRT3847_117455C.1.pap
US-10-424-599-162168

Query Match      29.9%; Score 29; DB 15; Length 47;
Best Local Similarity 38.5%; Pred. No. 2.1e+03;
Matches 5; Conservative 5; Mismatches 3; Indels 0; Gaps 0;

QY      3 NHLNSKIAFKIVS 15
DB      30 NILKNQIIFKLIN 42

RESULT 135
US-10-599-166629
; Sequence 166629, Application US/10424599
; Publication No. US20040031072A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa Thomas J
; APPLICANT: Kovalic David K
; APPLICANT: Zhou Yihua
; APPLICANT: Cao Yongwei
; TITLE OF INVENTION: Soy Nucleic Acid Molecules and Other Molecules Associated With
; TITLE OF INVENTION: Plants and Uses Thereof for Plant Improvement
; FILE REFERENCE: 38-21(53223)B
; CURRENT APPLICATION NUMBER: US/10/424,599
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